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(54) Title: CAULIFLOWER FLORAL MERISTEM IDENTITY GENES AND METHODS OF USING SAME

(57) Abstract

The present invention provides a nucleic acid molecule encoding a CAULIFLOWER (CAL) gene product such as a nucleic acid molecule encoding *Arabidopsis thaliana* CAL and a nucleic acid molecule encoding *Brassica oleracea* CAL (BoCAL). The invention also provides a nucleic acid molecule encoding a truncated CAL gene product such as a nucleic acid molecule encoding *Brassica oleracea* var. *botrytis* CAL (BobCAL). The invention also provides a nucleic acid containing the *Arabidopsis thaliana* CAL gene, a nucleic acid molecule containing the *Brassica oleracea* CAL gene and a nucleic acid molecule containing the *Brassica oleracea* var. *botrytis* CAL gene. The invention further provides a kit for converting shoot meristem to floral meristem and a kit for promoting early flowering in an angiosperm. The invention provides a CAL polypeptide and an antibody that specifically binds CAL polypeptides. In addition, the invention provides a truncated BobCAL polypeptide and an antibody that specifically binds truncated BobCAL polypeptide. The invention further provides a method of identifying a *Brassica* having a modified CAL allele by detecting a polymorphism associated with a CAL CAL locus, where the CAL CAL locus comprises a modified CAL CAL allele that does not encode an active CAL gene product.

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**CAULIFLOWER FLORAL MERISTEM IDENTITY GENES
AND METHODS OF USING SAME**

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BACKGROUND OF THE INVENTION

FIELD OF THE INVENTION

This invention relates generally to the field of plant flowering and more specifically to genes involved in the regulation of flowering.

BACKGROUND INFORMATION

A flower is the reproductive structure of a flowering plant. Following fertilization, the ovary of the flower becomes a fruit and bears seeds. As a practical consequence, production of fruit and seed-derived crops such as grapes, beans, corn, wheat and rice is dependent upon flowering.

Early in the plant life cycle, vegetative growth occurs, and roots, stems and leaves are formed. During the later period of reproductive growth, flowers as well as new shoots or branches develop. However, the factors responsible for the transition from vegetative to reproductive growth, and the onset of flowering, are poorly understood.

A variety of external signals, such as length of daylight and temperature, affect the time of flowering. The time of flowering also is subject to genetic controls that prevent young plants from flowering 5 prematurely. Thus, the pattern of genes expressed in a plant is an important determinant of the time of flowering.

Given these external signals and genetic controls, a relatively fixed period of vegetative growth 10 precedes flowering in a particular plant species. The length of time required for a crop to mature to flowering limits the geographic location in which it can be grown and can be an important determinant of yield. In addition, since the time of flowering determines when a 15 plant is reproductively mature, the pace of a plant breeding program also depends upon the length of time required for a plant to flower.

Traditionally, plant breeding involves generating hybrids of existing plants, which are examined 20 for improved yield or quality. The improvement of existing plant crops through plant breeding is central to increasing the amount of food grown in the world since the amount of land suitable for agriculture is limited. For example, the development of new strains of wheat, 25 corn and rice through plant breeding has increased the yield of these crops grown in underdeveloped countries such as Mexico, India and Pakistan. Unfortunately, plant breeding is inherently a slow process since plants must

be reproductively mature before selective breeding can proceed.

For some plant species, the length of time needed to mature to flowering is so long that selective breeding, which requires several rounds of backcrossing progeny plants with their parents, is impractical. For example, perennial trees such as walnut, hickory, oak, maple and cherry do not flower for several years after planting. As a result, breeding of such plant species for insect or disease-resistance or to produce improved wood or fruit, for example, would require many years, even if only a few rounds of selection were performed.

Methods of promoting early flowering can make breeding of long generation plants such as trees practical for the first time. Methods of promoting early flowering also would be useful for shortening growth periods, thereby broadening the geographic range in which a crop such as rice, corn or coffee can be grown. Unfortunately, methods for promoting early flowering in a plant have not yet been described. Thus, there is a need for methods that promote early flowering. The present invention satisfies this need and provides related advantages as well.

SUMMARY OF THE INVENTION

The present invention provides a nucleic acid molecule encoding a CAULIFLOWER (CAL) gene product. For example, the invention provides a nucleic acid molecule 5 encoding *Arabidopsis thaliana* CAL and a nucleic acid molecule encoding *Brassica oleracea* CAL.

The invention also provides a nucleic acid molecule encoding a truncated CAL gene product. For example, the invention provides a nucleic acid molecule 10 encoding the truncated *Brassica oleracea* var. *botrytis* CAL gene product. The invention also provides a nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule encoding a CAL gene product, a truncated CAL gene product, or a 15 complementary sequence thereto.

The invention further provides the *Arabidopsis thaliana* CAL gene, *Brassica oleracea* CAL gene and *Brassica oleracea* var. *botrytis* CAL gene. In addition, the invention provides a nucleotide sequence that 20 hybridizes under relatively stringent conditions to the *Arabidopsis thaliana* CAL gene, *Brassica oleracea* CAL gene or *Brassica oleracea* var. *botrytis* CAL gene, or a complementary sequence thereto.

The invention also provides vectors, including expression vectors, containing a nucleic acid molecule encoding a CAL gene product. The invention further provides a kit for converting shoot meristem to floral 5 meristem in an angiosperm and a kit for promoting early flowering in an angiosperm.

In addition, the invention provides a CAL polypeptide, such as the *Arabidopsis thaliana* CAL polypeptide or the *Brassica oleracea* CAL polypeptide, as 10 well as an antibody that specifically binds a CAL polypeptide. The invention further provides the truncated *Brassica oleracea* var. *botrytis* CAL polypeptide and an antibody that specifically binds the truncated *Brassica oleracea* var. *botrytis* CAL polypeptide.

15 The invention further provides a method of identifying a *Brassica* having a modified CAL allele by detecting a polymorphism associated with a CAL locus, where the CAL locus comprises a modified CAL allele that does not encode an active CAL gene product. For example, 20 the polymorphism can be a restriction fragment length polymorphism and the modified CAL allele can be the *Brassica oleracea* var. *botrytis* CAL allele.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 illustrates the nucleotide (SEQ ID 25 NO: 1) and amino acid (SEQ ID NO: 2) sequence of the *Arabidopsis thaliana* AP1 cDNA.

Figure 2 illustrates the nucleotide (SEQ ID NO: 3) and amino acid (SEQ ID NO: 4) sequence of the *Brassica oleracea* API cDNA.

Figure 3 illustrates the nucleotide (SEQ ID NO: 5) and amino acid (SEQ ID NO: 6) sequence of the *Brassica oleracea* var. *botrytis* API cDNA.

Figure 4 illustrates the nucleotide (SEQ ID NO: 7) and amino acid (SEQ ID NO: 8) sequence of the *Zea mays* API cDNA. The GenBank accession number is L46400.

10 Figure 5 illustrates the nucleotide (SEQ ID NO: 9) and amino acid (SEQ ID NO: 10) sequence of the *Arabidopsis thaliana* CAL cDNA.

15 Figure 6 illustrates the nucleotide (SEQ ID NO: 11) and amino acid (SEQ ID NO: 12) sequence of the *Brassica oleracea* CAL cDNA.

Figure 7 illustrates the nucleotide (SEQ ID NO: 13) and amino acid (SEQ ID NO: 14) sequence of the *Brassica oleracea* var. *botrytis* CAL cDNA.

20 Figure 8 illustrates CAL gene structure and provides a comparison of various CAL amino acid sequences.

Figure 8A. Exon-intron structure of *Arabidopsis* CAL gene. Exons are shown as boxes and introns as a solid line. Sizes (in base pairs) are

indicated above. Locations of changes resulting in mutant alleles are indicated by arrows. MADS and K domains are hatched.

Figure 8B. An alignment of three deduced amino acid sequences of CAL cDNAs. The complete *Arabidopsis thaliana* CAL amino acid sequence is displayed. The *Brassica oleracea* CAL (*BoCAL*) and *Brassica oleracea* var. *botrytis* CAL (*BobCAL*) amino acid sequences are shown directly below the *Arabidopsis* sequence where the sequences differ. The API amino acid sequence is shown for comparison. The MADS domain is indicated in bold and the K domain is underlined. GenBank accession numbers are as follows: *Arabidopsis thaliana* CAL (L36925); *Brassica oleracea* CAL (L36926) and *Brassica oleracea* var. *botrytis* CAL (L36927).

Figure 9 illustrates the nucleotide (SEQ ID NO: 15) and amino acid (SEQ ID NO: 16) sequence of the *Arabidopsis thaliana* LEAFY (LFY) cDNA.

Figure 10 illustrates the genomic sequence of *Arabidopsis thaliana* API (SEQ ID NO: 17).

Figure 11 illustrates the genomic sequence of *Brassica oleracea* API (SEQ ID NO: 18).

Figure 12 illustrates the genomic sequence of *Brassica oleracea* var. *botrytis* API (SEQ ID NO: 19).

Figure 13 illustrates the genomic sequence of *Arabidopsis thaliana* CAL (SEQ ID NO: 20).

Figure 14 illustrates the genomic sequence of *Brassica oleracea* CAL (SEQ ID NO: 21).

5 Figure 15 illustrates the genomic sequence of *Brassica oleracea* var. *botrytis* CAL (SEQ ID NO: 22).

Figure 16 illustrates the nucleotide (SEQ ID NO: 23) and amino acid (SEQ ID NO: 24) sequence of the rat glucocorticoid receptor ligand binding domain.

10

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides a nucleic acid molecule encoding a CAULIFLOWER (CAL) gene product, which is a floral meristem identity gene product involved in the conversion of shoot meristem to floral meristem. For 15 example, the invention provides a nucleic acid molecule encoding *Arabidopsis thaliana* CAL and a nucleic acid molecule encoding *Brassica oleracea* CAL (BoCAL) (Kempin et al., *Science*, 267:522-525 (1995), which is incorporated herein by reference). As disclosed herein, 20 a CAL gene product can be expressed in an angiosperm, thereby converting shoot meristem to floral meristem in the angiosperm or promoting early flowering in the angiosperm. The invention also provides a nucleic acid molecule encoding a truncated CAL gene product such as a 25 nucleic acid molecule encoding *Brassica oleracea* var. *botrytis* CAL (BobCAL). The invention also provides a

nucleic acid molecule containing the *Arabidopsis thaliana* CAL gene, a nucleic acid molecule containing the *Brassica oleracea* CAL gene and a nucleic acid molecule containing the *Brassica oleracea* var. *botrytis* CAL gene. The 5 invention further provides a kit for converting shoot meristem to floral meristem and a kit for promoting early flowering in an angiosperm. The invention provides a CAL polypeptide and an antibody that specifically binds CAL polypeptide. In addition, the invention provides the 10 truncated BobCAL polypeptide and an antibody that specifically binds the truncated BobCAL polypeptide. The invention further provides a method of identifying a *Brassica* having a modified CAL allele by detecting a polymorphism associated with a CAL locus, where the CAL 15 locus comprises a modified CAL allele that does not encode an active CAL gene product.

The present invention provides a non-naturally occurring angiosperm containing a first ectopically expressible nucleic acid molecule encoding a first floral 20 meristem identity gene product. For example, the invention provides a transgenic angiosperm containing a first ectopically expressible floral meristem identity gene product such as APETALA1 (AP1), CAULIFLOWER (CAL) or LEAFY (LFY). Such a transgenic angiosperm can be, for 25 example, a cereal plant, leguminous plant, oilseed plant, tree, fruit-bearing plant or ornamental flower.

A flower, like a leaf or shoot, is derived from the shoot apical meristem, which is a collection of 30 undifferentiated cells set aside during embryogenesis.

The production of vegetative structures, such as leaves or shoots, and of reproductive structures, such as flowers, is temporally segregated, such that a leaf or shoot arises early in a plant life cycle, while a flower 5 develops later. The transition from vegetative to reproductive development is the consequence of a process termed floral induction (Yanofsky, Ann. Rev. Plant Physiol. Plant Mol. Biol. 46:167-188 (1995)).

Once induced, shoot apical meristem either 10 persists and produces floral meristem, which gives rise to flowers, and lateral meristem, which gives rise to branches, or is itself converted to floral meristem. The fate of floral meristem is to differentiate into a single flower having a fixed number of floral organs in a 15 whorled arrangement. Dicots, for example, contain four whorls (concentric rings) in which sepals (first whorl) and petals (second whorl) surround stamens (third whorl) and carpels (fourth whorl).

Although shoot meristem and floral meristem 20 both consist of meristematic tissue, shoot meristem is distinguishable from the more specialized floral meristem. Shoot meristem generally is indeterminate and gives rise to an unspecified number of floral and lateral meristems. In contrast, floral meristem is determinate 25 and gives rise to the fixed number of floral organs that comprise a flower.

By convention herein, a wild-type gene sequence is represented in upper case italic letters (for example,

APETALA1), and a wild-type gene product is represented in upper case non-italic letters (*APETALA1*). Further, a mutant gene allele is represented in lower case italic letters (*apl*), and a mutant gene product is represented 5 in lower case non-italic letters (*apl*).

Genetic studies have identified a number of genes involved in regulating flower development. These genes can be classified into different groups depending on their function. Flowering time genes, for example, 10 are involved in floral induction and regulate the transition from vegetative to reproductive growth. In comparison, the floral meristem identity genes, which are the subject matter of the present invention as disclosed herein, encode proteins that promote the conversion of 15 shoot meristem to floral meristem. In addition, floral organ identity genes encode proteins that determine whether sepals, petals, stamens or carpels are formed (Yanofsky, *supra*, 1995; Weigel, *Ann. Rev. Genetics* 29:19-39 (1995)). Some of the floral meristem identity 20 gene products also have a role in specifying organ identity.

Floral meristem identity genes have been identified by characterizing genetic mutations that prevent or alter floral meristem formation. Among floral 25 meristem identity gene mutations in *Arabidopsis thaliana*, those in the gene *LEAFY* (*LFY*) generally have the strongest effect on floral meristem identity. Mutations in *LFY* completely transform the basal-most flowers into secondary shoots and have variable effects on

later-arising (apical) flowers. In comparison, mutations in the floral meristem identity gene *APETALA1* (*API*) result in replacement of a few basal flowers by inflorescence shoots that are not subtended by leaves.

5 An apical flower produced in an *api* mutant has an indeterminate structure in which a flower arises within a flower. These mutant phenotypes indicate that both *API* and *LFY* contribute to establishing the identity of the floral meristem although neither gene is absolutely required. The phenotype of *lfy api* double mutants, in which structures with flower-like characteristics are very rare, indicates that *LFY* and *API* encode partially redundant activities.

In addition to the *LFY* and *API* genes, a third locus that greatly enhances the *api* mutant phenotype has been identified in *Arabidopsis*. This locus, designated *CAULIFLOWER* (*CAL*), derives its name from the resulting "cauliflower" phenotype, which is strikingly similar to the common garden variety of cauliflower. In an *api cal* double mutant, floral meristem that develops behaves as shoot meristem in that there is a massive proliferation of meristems in the position that normally would be occupied by a single flower. However, a plant homozygous for a particular *cal* mutation (*cal-1*) has a normal phenotype, indicating that *API* can substitute for the loss of *CAL* in these plants. In addition, because floral meristem that forms in an *api* mutant behaves as shoot meristem in an *api cal* double mutant, *CAL* can largely substitute for *API* in specifying floral meristem. These 30 genetic data indicate that *CAL* and *API* encode activities

that are partially redundant in converting shoot meristem to floral meristem.

Other genetic loci play at least minor roles in specifying floral meristem identity. For example, 5 although a mutation in *APETALA2* (*AP2*) alone does not result in altered inflorescence characteristics, *ap2 api* double mutants have indeterminate flowers (flowers with shoot-like characteristics) (Bowman et al., *Development* 119:721-743 (1993)). Also, mutations in the *CLAVATA1* 10 gene result in an enlarged meristem and lead to a variety of phenotypes (Clark et al., *Development* 119:397-418 (1993)). In a *clv1 api* double mutant, formation of flowers is initiated, but the center of each flower often develops as an indeterminate inflorescence. 15 Thus, mutations in *CLAVATA1* result in the loss of floral meristem identity in the center of wild-type flowers. Genetic evidence also indicates that the gene product of *UNUSUAL FLORAL ORGANS* (*UFO*) plays a role in determining the identity of floral meristem. Additional floral 20 meristem identity genes associated with altered floral meristem formation remain to be isolated.

Mutations in another locus, designated *TERMINAL FLOWER* (*TFL*), produce phenotypes that generally are reversed as compared to mutations in the floral meristem 25 identity genes. For example, *tfl* mutants flower early, and the indeterminate apical and lateral meristems develop as determinate floral meristems (Alvarez et al., *Plant J.* 2:103-116 (1992)). These characteristics indicate that the *TFL* promotes maintenance of shoot

meristem. TFL also acts directly or indirectly to negatively regulate AP1 and LFY expression in shoot meristem since AP1 and LFY are ectopically expressed in the shoot meristem of *tfl* mutants (Gustafson-Brown et al., *Cell* 76:131-143 (1994); Weigel et al., *Cell* 69:843-859 (1992)). It is recognized that a plant having a mutation in TFL can have a phenotype similar to a non-naturally occurring angiosperm of the invention. Such *tfl* mutants, however, are explicitly excluded from the scope of the present invention.

The results of such genetic studies indicate that several floral meristem identity gene products, including AP1, CAL and LFY, act redundantly to convert shoot meristem to floral meristem and that TFL acts directly or indirectly to negatively regulate expression of the floral meristem identity genes. As disclosed herein, ectopic expression of a single floral meristem identity gene product such as AP1, CAL or LFY is sufficient to convert shoot meristem to floral meristem. Thus, the present invention provides a non-naturally occurring angiosperm that contains an ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product, provided that such ectopic expression is not due to a mutation in an endogenous TERMINAL FLOWER gene.

As disclosed herein, an ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product can be, for example, a transgene encoding a floral meristem identity gene product under control of a

heterologous gene regulatory element. In addition, such an ectopically expressible nucleic acid molecule can be an endogenous floral meristem identity gene coding sequence that is placed under control of a heterologous 5 gene regulatory element. The ectopically expressible nucleic acid molecule also can be, for example, an endogenous floral meristem identity gene having a modified gene regulatory element such that the endogenous floral meristem identity gene is no longer subject to 10 negative regulation by TFL.

The term "ectopically expressible" is used herein to refer to a gene transcript or gene product that can be expressed in a tissue other than a tissue in which it normally is produced. The actual ectopic expression 15 thereof is dependent on various factors and can be constitutive or inducible expression. As disclosed herein, AP1, which normally is expressed in floral meristem, is ectopically expressible in shoot meristem. As disclosed herein, when a floral meristem identity gene 20 product such as AP1, CAL or LFY is ectopically expressed in shoot meristem, the shoot meristem is converted to floral meristem and early flowering can occur (see Examples II, IV and V).

In particular, an ectopically expressible 25 nucleic acid molecule encoding a floral meristem identity gene product can be expressed prior to the developmental time at which the corresponding endogenous gene normally is expressed. For example, an *Arabidopsis* plant grown under continuous light conditions expresses AP1 just

prior to day 18, when normal flowering begins. However, as disclosed herein, *AP1* can be ectopically expressed in shoot meristem earlier than day 18, resulting in early conversion of shoot meristem to floral meristem and early 5 flowering. As shown in Example IID, a transgenic *Arabidopsis* plant that ectopically expresses *AP1* in shoot meristem under control of a constitutive promoter flowers earlier than the corresponding non-transgenic plant (day 10 as compared to day 18).

10 As used herein, the term "floral meristem identity gene product" means a gene product that promotes conversion of shoot meristem to floral meristem. As disclosed herein, expression of a floral meristem identity gene product such as *AP1*, *CAL* or *LFY* in shoot 15 meristem can convert shoot meristem to floral meristem. Furthermore, expression of a floral meristem identity gene product in shoot meristem also can promote early flowering (Examples IID, IVA and V). A floral meristem identity gene product is distinguishable from a late 20 flowering gene product or an early flowering gene product, which are not encompassed within the present invention. In addition, reference is made herein to an "inactive" floral meristem identity gene product, as exemplified by *BobCAL* (see below). Expression of an 25 inactive floral meristem identity gene product in an angiosperm does not result in the conversion of shoot meristem to floral meristem in the angiosperm.

A floral meristem identity gene product can be, for example, an *AP1* gene product such as *Arabidopsis AP1*,

which is a 256 amino acid gene product encoded by the AP1 cDNA sequence isolated from *Arabidopsis thaliana* (Figure 5, SEQ ID NO: 2). The *Arabidopsis* AP1 cDNA encodes a highly conserved MADS domain, which can 5 function as a DNA-binding domain, and a K domain, which is structurally similar to the coiled-coil domain of keratins and can be involved in protein-protein interactions.

In *Arabidopsis*, AP1 RNA is expressed in flowers 10 but is not detectable in roots, stems or leaves (Mandel et al., *Nature* 360:273-277 (1992), which is incorporated herein by reference). The earliest detectable expression of AP1 RNA is in young floral meristem at the time it initially forms on the flanks of shoot meristem. 15 Expression of AP1 increases as the floral meristem increases in size; no AP1 expression is detectable in shoot meristem. In later stages of development, AP1 expression ceases in cells that will give rise to reproductive organs (stamens and carpels), but is 20 maintained in cells that will give rise to non-reproductive organs (sepals and petals; Mandel, *supra*, 1992).

As used herein, the term "APETALA1" or "AP1" means a floral meristem identity gene product that is 25 characterized, in part, by having an amino acid sequence that is related to the *Arabidopsis* AP1 amino acid sequence shown in Figure 1 (SEQ ID NO: 2) or to the *Zea mays* AP1 amino acid sequence shown in Figure 4 (SEQ ID NO: 8). In nature, AP1 is expressed in floral meristem.

CAULIFLOWER (CAL) is another example of a floral meristem identity gene product. As used herein, the term "CAULIFLOWER" or "CAL" means a floral meristem identity gene product that is characterized in part by

5 having an amino acid sequence that has at least about 70 percent identity with the amino acid sequence shown in Figure 5 (SEQ ID NO: 10) in the region from amino acid 1 to amino acid 160 or with the amino acid sequence shown in Figure 6 (SEQ ID NO: 12) in the region from amino acid

10 1 to amino acid 160. In nature, CAL is expressed in floral meristem.

The present invention provides a nucleic acid molecule encoding a CAL, including, for example, the *Arabidopsis* CAL cDNA sequence shown in Figure 5 (SEQ ID NO: 9). As disclosed herein, CAL, like AP1, contains a MADS domain and a K domain. The MADS domains of CAL and AP1 differ in only five of 56 amino acid residues, where four of the five differences represent conservative amino acid replacements. Over the entire sequence, the

15 *Arabidopsis* CAL and *Arabidopsis* AP1 sequences (SEQ ID NOS: 10 and 2) are 76% identical and are 88% similar if conservative amino acid substitutions are allowed.

Similar to the expression pattern of AP1, CAL RNA is expressed in young floral meristem in *Arabidopsis*.

20 However, in contrast to AP1 expression, which is high throughout sepal and petal development, CAL expression is low in these organs.

LEAFY (LFY) is yet another example of a floral meristem identity gene product. As used herein, the term "LEAFY" or "LFY" means a floral meristem identity gene product that is characterized in part by having an amino acid sequence that is related to the amino acid sequence shown in Figure 9 (SEQ ID NO: 16). In nature, LFY is expressed in floral meristem as well as during vegetative development. As disclosed herein, ectopic expression of floral meristem identity gene products, which normally 5 are expressed in floral meristem, such as AP1 or CAL or LFY or combinations thereof, in shoot meristem can 10 convert shoot meristem to floral meristem and promote early flowering.

Flower development in *Arabidopsis* is recognized 15 in the art as a model for flower development in angiosperms in general. Gene orthologs corresponding to the *Arabidopsis* genes involved in the early steps of flower formation have been identified in distantly related plant species, and these gene orthologs show 20 remarkably similar RNA expression patterns. Mutations in these genes also result in phenotypes that correspond to the phenotype produced by a similar mutation in *Arabidopsis*. For example, orthologs of the *Arabidopsis* floral meristem identity genes *AP1* and *LFY* and the 25 *Arabidopsis* organ identity genes *AGAMOUS*, *APETALA3* and *PISTILLATA* have been isolated from monocots such as maize and, where characterized, reveal the anticipated RNA expression patterns and related mutant phenotypes. (Schmidt et al., Plant Cell 5:729-737 (1993); and Veit et 30 al., Plant Cell 5:1205-1215 (1993), each of which is

incorporated herein by reference). Furthermore, a gene ortholog can be functionally interchangeable in that it can function across distantly related species boundaries (Mandel et al., Cell 71:133-143 (1992), which is incorporated herein by reference). Taken together, these data suggest that the underlying mechanisms controlling the initiation and proper development of flowers are conserved across distantly related dicot and monocot boundaries. Therefore, results obtained using *Arabidopsis* can be predictive of results that can be expected in other angiosperms.

Floral meristem identity genes in particular are conserved throughout the plant kingdom. For example, a gene ortholog of *Arabidopsis API* has been isolated from *Antirrhinum majus* (snapdragon; Huijser et al., EMBO J. 11:1239-1249 (1992), which is herein incorporated by reference). As disclosed herein, an ortholog of *Arabidopsis API* also has been isolated from *Zea Mays* (maize; see Example IA). Similarly, gene orthologs of *Arabidopsis LFY* have been isolated from *Antirrhinum majus*, tobacco and poplar tree (Coen et al., Cell, 63:1311-1322 (1990); Kelly et al., Plant Cell 7:225-234 (1995); and Strauss et al., Molec. Breed 1:5-26 (1995), each of which is incorporated herein by reference). In addition, a mutation in the *Antirrhinum API* ortholog results in a phenotype similar to the *Arabidopsis apl* mutant phenotype described above (Huijser et al., *supra*, 1992). Similarly, a mutation in the *Antirrhinum LFY* ortholog results in a phenotype similar to the *Arabidopsis lfy* mutant phenotype (Coen et al., *supra*,

1995). These studies indicate that *AP1* and *LFY* function similarly in distantly related angiosperms.

A floral meristem identity gene product also can function across species boundaries. For example,
5 *Arabidopsis LFY* can convert shoot meristem to floral meristem when expressed in aspen trees (Weigel and Nilsson, *Nature* 377:495-500 (1995), which is incorporated herein by reference). As disclosed herein, a nucleic acid molecule encoding an *Arabidopsis AP1* or *CAL* gene
10 product (SEQ ID NOS: 1 and 9), for example, also can be used to convert shoot meristem to floral meristem in an angiosperm. Thus, a nucleic acid molecule encoding an *Arabidopsis AP1* gene product (SEQ ID NO: 1) or an *Arabidopsis CAL* gene product (SEQ ID NO: 9) can be
15 introduced into an angiosperm such as corn, wheat or rice and, upon expression, can convert shoot meristem to floral meristem in the transgenic angiosperm. Furthermore, as disclosed herein, the conserved nature of an *AP1* or *CAL* or *LFY* gene among diverse angiosperms,
20 allows a nucleic acid molecule encoding a floral meristem identity gene product from essentially any angiosperm to be introduced into essentially any other angiosperm, wherein the expression of the nucleic acid molecule in shoot meristem can convert shoot meristem to floral
25 meristem.

If desired, a novel *AP1*, *CAL* or *LFY* sequence can be isolated from an angiosperm using a nucleotide sequence as a probe and methods well known in the art of molecular biology (Sambrook et al. (eds.), *Molecular*

Cloning: A Laboratory Manual (Second Edition),
Plainview, NY: Cold Spring Harbor Laboratory Press
(1989), which is herein incorporated by reference). As
exemplified herein and discussed in detail below (see
5 Example IA), the *AP1* ortholog from *Zea Mays* (maize; SEQ
ID NO: 7) was isolated using the *Arabidopsis AP1* cDNA as
a probe (SEQ ID NO: 1).

In one embodiment, the invention provides a
non-naturally occurring angiosperm that contains an
10 ectopically expressible nucleic acid molecule encoding a
floral meristem identity gene product and that is
characterized by early flowering. As used herein, the
term "characterized by early flowering," when used in
reference to a non-naturally occurring angiosperm of the
15 invention, means a non-naturally occurring angiosperm
that forms flowers sooner than flowers would form on a
corresponding naturally occurring angiosperm that does
not ectopically express a floral meristem identity gene
product, grown under the same conditions. Flowering
20 times for naturally occurring angiosperms are well known
in the art and depend, in part, on genetic factors and on
the environmental conditions, such as day length. Thus,
given a defined set of environmental conditions, a
naturally occurring plant will flower at a relatively
25 predictable time.

It is recognized that various transgenic plants
that are characterized by early flowering have been
described. Such transgenic plants are described herein
and are readily distinguishable or explicitly excluded

from the present invention. For example, a product of a "late-flowering gene" can promote early flowering but does not specify the conversion of shoot meristem to floral meristem. Therefore, a transgenic plant
5 expressing a late-flowering gene product is distinguishable from a non-naturally occurring angiosperm of the invention. For example, a transgenic plant expressing the late-flowering gene, *CONSTANS* (*CO*), flowers earlier than a corresponding wild type plant
10 (Putterill et al., *Cell* 80:847-857 (1995)). However, expression of exogenous *CONSTANS* does not convert shoot meristem to floral meristem.

Early flowering also has been observed in a transgenic tobacco plant expressing an exogenous rice
15 MADS domain gene. Although the product of this gene promotes early flowering, it does not specify the identity of floral meristem and, thus, cannot convert shoot meristem to floral meristem (Chung et al., *Plant Mol. Biol.* 26:657-665 (1994)). Therefore, the
20 early-flowering CO and rice MADS domain gene transgenic plants are distinguishable from the early-flowering non-naturally occurring angiosperms of the invention.

Mutations in a class of genes known as "early-flowering genes" also result in plants that flower
25 prematurely. Such early flowering genes include, for example, *EARLY FLOWERING 1-3* (*ELF1*, *ELF2*, *ELF3*); *EMBRYONIC FLOWER 1,2* (*EMF1*, *EMF2*); *LONG HYPOCOTYL 1,2* (*HY1*, *HY2*); *PHYTOCHROME B* (*PHYB*), *SPINDLY* (*SPY*) and *TERMINAL FLOWER* (*TFL*) (Weigel, *supra*, 1995). However,

the wild type product of an early flowering gene retards flowering and is distinguishable from a floral meristem identity gene product in that it does not promote conversion of shoot meristem to floral meristem.

5 An *Arabidopsis* plant having a mutation in the *TERMINAL FLOWER (TFL)* gene flowers early and is characterized by the conversion of shoots to flowers (Alvarez et al., *Plant J.* 2:103-116 (1992), which is incorporated herein by reference). However, TFL is not a
10 floral meristem identity gene product, as defined herein. Specifically, it is the loss of TFL that promotes conversion of shoot meristem to floral meristem. Since the function of TFL is to antagonize formation of floral meristem, a *tfl* mutant, which has lost this antagonist
15 function, permits conversion of shoot meristem to floral meristem. Although TFL is not a floral meristem identity gene product and does not itself convert shoot meristem to floral meristem, the loss of TFL can result in a plant with an ectopically expressed floral meristem identity
20 gene product. Such *tfl* mutants, in which a mutation in *TFL* results in conversion of shoot meristem to floral meristem, are explicitly excluded from the present invention.

As used herein, the term "non-naturally occurring angiosperm" means an angiosperm that contains a genome that has been modified by man. A transgenic angiosperm, for example, contains an exogenous nucleic acid molecule and, therefore, contains a genome that has been modified by man. Furthermore, an angiosperm that

contains, for example, a mutation in an endogenous floral meristem identity gene regulatory element as a result of exposure to a mutagenic agent by man also contains a genome that has been modified by man. In contrast, a 5 plant containing a spontaneous or naturally occurring mutation is not a "non-naturally occurring angiosperm" and, therefore, is not encompassed within the invention.

As used herein, the term "transgenic" refers to an angiosperm that contains in its genome an exogenous 10 nucleic acid molecule, which can be derived from the same or a different species. The exogenous nucleic acid molecule that is introduced into the angiosperm can be a gene regulatory element such as a promoter or other regulatory element or can be a coding sequence, which can 15 be linked to a heterologous gene regulatory element.

As used herein, the term "angiosperm" means a flowering plant. Angiosperms are well known and produce a variety of useful products including materials such as lumber, rubber, and paper; fibers such as cotton and 20 linen; herbs and medicines such as quinine and vinblastine; ornamental flowers such as roses and orchids; and foodstuffs such as grains, oils, fruits and vegetables.

Angiosperms are divided into two broad classes 25 based on the number of cotyledons, which are seed leaves that generally store or absorb food. Thus, a monocotyledonous angiosperm is an angiosperm having a

single cotyledon, and a dicotyledonous angiosperm is an angiosperm having two cotyledons.

Angiosperms encompass a variety of flowering plants, including, for example, cereal plants, leguminous plants, oilseed plants, trees, fruit-bearing plants and ornamental flowers, which general classes are not necessarily exclusive. Such angiosperms include for example, a cereal plant, which produces an edible grain cereal. Such cereal plants include, for example, corn, rice, wheat, barley, oat, rye, orchardgrass, guinea grass, sorghum and turfgrass. In addition, a leguminous plant is an angiosperm that is a member of the pea family (Fabaceae) and produces a characteristic fruit known as a legume. Examples of leguminous plants include, for example, soybean, pea, chickpea, moth bean, broad bean, kidney bean, lima bean, lentil, cowpea, dry bean, and peanut. Examples of legumes further also include alfalfa, birdsfoot trefoil, clover and sainfoin. Furthermore, an oilseed plant is an angiosperm that has seeds useful as a source of oil. Examples of oilseed plants include soybean, sunflower, rapeseed and cottonseed.

A tree is an angiosperm and is a perennial woody plant, generally with a single stem (trunk).
25 Examples of trees include alder, ash, aspen, basswood (linden), beech, birch, cherry, cottonwood, elm, eucalyptus, hickory, locust, maple, oak, persimmon, poplar, sycamore, walnut and willows. Such trees are

used for pulp, paper, and structural material, as well as providing a major source of fuel.

A fruit-bearing plant also is an angiosperm and produces a mature, ripened ovary (usually containing 5 seeds) that is suitable for human or animal consumption. Examples of fruit-bearing plants include grape, orange, lemon, grapefruit, avocado, date, peach, cherry, olive, plum, coconut, apple and pear trees and blackberry, blueberry, raspberry, strawberry, pineapple, tomato, 10 cucumber and eggplant plants. An ornamental flower is an angiosperm cultivated for its decorative flower. Examples of ornamental flowers include rose, orchid, lily, tulip and chrysanthemum, snapdragon, camelia, carnation and petunia. The skilled artisan will 15 recognize that the invention can be practiced on these or other angiosperms, as desired.

In various embodiments, the present invention provides a non-naturally occurring angiosperm having an ectopically expressible first nucleic acid molecule 20 encoding a first floral meristem identity gene product, provided the first nucleic acid molecule is not ectopically expressed due to a mutation in an endogenous TFL gene. If desired, a non-naturally occurring angiosperm of the invention can contain an ectopically 25 expressible second nucleic acid molecule encoding a second floral meristem identity gene product, which is different from the first floral meristem identity gene product.

An ectopically expressible nucleic acid molecule can be expressed, as desired, either constitutively or inducibly. Such an ectopically expressible nucleic acid molecule can be an endogenous 5 nucleic acid molecule and can contain, for example, a mutation in its endogenous gene regulatory element or can contain an exogenous, heterologous gene regulatory element that is linked to and directs expression of the endogenous nucleic acid molecule. In addition, an 10 ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product can be an exogenous nucleic acid molecule encoding a floral meristem identity gene product and containing a heterologous gene regulatory element.

15 The invention provides, for example, a non-naturally occurring angiosperm containing a first ectopically expressible nucleic acid molecule encoding a first floral meristem identity gene product. If desired, a non-naturally occurring angiosperm of the invention can 20 contain a floral meristem identity gene having a modified gene regulatory element and also can contain a second ectopically expressible nucleic acid molecule encoding a second floral meristem identity gene product, provided that neither the first nor second ectopically expressible 25 nucleic acid molecule is ectopically expressed due to a mutation in an endogenous TERMINAL FLOWER gene.

As used herein, the term "modified gene regulatory element" means a regulatory element having a mutation that results in ectopic expression in shoot

meristem of the floral meristem identity gene regulated by the gene regulatory element. Such a gene regulatory element can be, for example, a promoter or enhancer element and can be positioned 5' or 3' to the coding sequence or within an intronic sequence of the floral meristem identity gene. Such a modification can be, for example, a nucleotide insertion, deletion or substitution and can be produced by chemical mutagenesis using a mutagen such as ethylmethane sulfonate (see Example IIIA) or by insertional mutagenesis using a transposable element. For example, a modified gene regulatory element can be a functionally inactivated binding site for TFL or a gene product regulated by TFL, such that modification of the gene regulatory element results in ectopic expression of the floral meristem identity gene product in shoot meristem.

The invention also provides a transgenic angiosperm containing a first exogenous gene promoter that regulates a first ectopically expressible nucleic acid molecule encoding a first floral meristem identity gene product and a second exogenous gene promoter that regulates a second ectopically expressible nucleic acid molecule encoding a second floral meristem identity gene product.

The invention also provides a transgenic angiosperm containing a first exogenous ectopically expressible nucleic acid molecule encoding a first floral meristem identity gene product and a second exogenous gene promoter that regulates a second ectopically

expressible nucleic acid molecule encoding a second floral meristem identity gene product, provided that the first nucleic acid molecule is not ectopically expressed due to a mutation in an endogenous *TERMINAL FLOWER* gene.

5 The invention also provides a transgenic angiosperm containing a first exogenous ectopically expressible nucleic acid molecule encoding a first floral meristem identity gene product and a second exogenous ectopically expressible nucleic acid molecule encoding a
10 second floral meristem identity gene product, where the first floral meristem identity gene product is different from the second floral meristem identity gene product and provided that neither nucleic acid molecule is ectopically expressed due to a mutation in an endogenous
15 *TERMINAL FLOWER* gene.

The ectopic expression of first and second floral meristem identity gene products can be particularly useful. For example, ectopic expression of AP1 and LFY in a plant promotes flowering earlier than
20 ectopic expression of AP1 alone or ectopic expression of LFY alone. Thus, plant breeding, for example, can be further accelerated, if desired.

First and second floral meristem identity gene products can be, for example, AP1 and CAL, or can be AP1
25 and LFY or can be CAL and LFY. It should be recognized that where a transgenic angiosperm of the invention contains two exogenous nucleic acid molecules, the order of introducing such a first and a second nucleic acid

molecule is not important for purposes of the present invention. Thus, a transgenic angiosperm of the invention having, for example, AP1 as the first floral meristem identity gene product and CAL as the second 5 floral meristem identity gene product is equivalent to a transgenic angiosperm having CAL as the first floral meristem identity gene product and AP1 as the second floral meristem identity gene product.

The invention also provides methods of
10 converting shoot meristem to floral meristem in an angiosperm by ectopically expressing an ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product in the angiosperm. Thus, the invention provides, for example, methods of
15 converting shoot meristem to floral meristem in an angiosperm by introducing an exogenous ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product into the angiosperm, thereby producing a transgenic angiosperm. A floral
20 meristem identity gene product such as AP1, CAL or LFY, or a chimeric protein containing, in part, a floral meristem identity gene product (see below) is useful in the methods of the invention.

As used herein, the term "introducing," when
25 used in reference to an angiosperm, means transferring an exogenous nucleic acid molecule into the angiosperm. For example, an exogenous nucleic acid molecule can be introduced into an angiosperm by methods such as *Agrobacterium*-mediated transformation or direct gene

transfer methods including microprojectile-mediated transformation (Klein et al., Nature 327:70-73 (1987), which is incorporated herein by reference). These and other methods of introducing a nucleic acid molecule into 5 an angiosperm are well known in the art (Bowman et al. (ed.), Arabidopsis: An Atlas of Morphology and Development, New York: Springer (1994); Valvekens et al., Proc. Natl. Acad. Sci. USA 85:5536-5540 (1988); and Wang et al., Transformation of Plants and Soil 10 Microorganisms, Cambridge, UK: University Press (1995), each of which is incorporated herein by reference).

As used herein, the term "converting shoot meristem to floral meristem" means promoting the formation of flower progenitor tissue where shoot 15 progenitor tissue would normally be formed. As a result of the conversion of shoot meristem to floral meristem, flowers form in an angiosperm where shoots normally would form. The conversion of shoot meristem to floral meristem can be identified using well known methods, such 20 as scanning electron microscopy, light microscopy or visual inspection.

The invention also provides methods of converting shoot meristem to floral meristem in an angiosperm by introducing a first ectopically expressible 25 nucleic acid molecule encoding a first floral meristem identity gene product and a second ectopically expressible nucleic acid molecule encoding a second floral meristem identity gene product into the angiosperm. As discussed above, first and second floral

meristem identity gene products useful in the invention can be, for example, AP1 and CAL or AP1 and LFY or CAL and LFY.

- The invention also provides methods of
- 5 promoting early flowering in an angiosperm by ectopically expressing a nucleic acid molecule encoding a floral meristem identity gene product in the angiosperm, provided that the nucleic acid molecule is not ectopically expressed due to a mutation in an endogenous
- 10 *TERMINAL FLOWER* gene. For example, the invention provides methods of promoting early flowering in an angiosperm by introducing an ectopically expressible nucleic acid molecule encoding a floral meristem identity gene product into the angiosperm, thus producing a
- 15 transgenic angiosperm. A floral meristem identity gene product such as AP1, CAL or LFY, or a chimeric protein containing, in part, a floral meristem identity gene product (see below) is useful in methods of promoting early flowering.
- 20 The present invention further provides nucleic acid molecules encoding floral meristem identity gene products. For example, the invention provides a nucleic acid molecule encoding CAL, having at least about 70 percent amino acid identity with amino acids 1 to 160 of
- 25 SEQ ID NO: 10 or SEQ ID NO: 11. The invention also provides a nucleic acid molecule encoding *Arabidopsis thaliana* CAL having the amino acid sequence shown in Figure 5 (SEQ ID NO: 10) and a nucleic acid molecule encoding *Brassica oleracea* CAL having the amino acid

sequence shown in Figure 6 (SEQ ID NO: 12). In addition, the invention provides a nucleic acid molecule encoding *Brassica oleracea* AP1 having the amino acid sequence shown in Figure 2 (SEQ ID NO: 4) and a nucleic acid 5 molecule encoding *Brassica oleracea* var. *botrytis* AP1 having the amino acid sequence shown in Figure 3 (SEQ ID NO: 6). The invention also provides a nucleic acid molecule encoding *Zea mays* AP1 having the amino acid sequence shown in Figure 4 (SEQ ID NO: 8).

10 As disclosed herein, CAL is highly conserved among different angiosperms. For example, *Arabidopsis* CAL (SEQ ID NO: 10) and *Brassica oleracea* CAL (SEQ ID NO: 12) share about 80 percent amino acid identity. In the region from amino acid 1 to amino acid 160, *Arabidopsis* 15 CAL and *Brassica oleracea* CAL are about 89 percent identical at the amino acid level. Using a nucleotide sequence derived from a conserved region of SEQ ID NO: 9 or SEQ ID NO: 11, a nucleic acid molecule encoding a novel CAL ortholog can be isolated from other 20 angiosperms. Using methods such as those described by Purugganan et al. (*Genetics* 40: 345-356 (1995)), one can readily confirm that the newly isolated molecule is a CAL ortholog. Thus, a nucleic acid molecule encoding CAL, which has at least about 70 percent amino acid identity 25 with *Arabidopsis* CAL (SEQ ID NO: 10) or *Brassica oleracea* CAL (SEQ ID NO: 12), can be isolated and identified using well known methods.

The invention also provides a nucleic acid molecule encoding a truncated CAL gene product. For

example, the invention provides a nucleic acid molecule encoding the *Brassica oleracea* var. *botrytis* CAL gene product (BobCAL). BobCAL contains 150 amino acids of the approximately 255 amino acids encoded by a full-length 5 CAL cDNA (see Figure 7; SEQ ID NO: 14; see, also, Figure 8B).

The invention also provides a nucleic acid containing the *Arabidopsis thaliana* API gene (Figure 10; SEQ ID NO: 17), a nucleic acid molecule containing the 10 *Brassica oleracea* API gene (Figure 11; SEQ ID NO: 18) and a nucleic acid molecule containing the *Brassica oleracea* var. *botrytis* API gene (Figure 12; SEQ ID NO: 19). In addition, the invention also provides a nucleic acid containing the *Arabidopsis thaliana* CAL gene (Figure 13; 15 SEQ ID NO: 20) and a nucleic acid molecule containing the *Brassica oleracea* CAL gene (Figure 11; SEQ ID NO: 21). In addition, the invention provides a nucleic acid molecule containing the *Brassica oleracea* var. *botrytis* CAL gene (Figure 15; SEQ ID NO: 22).

20

The invention further provides a nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule encoding a CAL, or a complementary sequence thereof. In particular, such a 25 nucleotide sequence can hybridize under relatively stringent conditions to a nucleic acid molecule encoding *Arabidopsis* CAL (SEQ ID NO: 9) or *Brassica oleracea* CAL (SEQ ID NO: 11), or a complementary sequence thereof. Similarly, the present invention provides a nucleotide 30 sequence that hybridizes under relatively stringent

conditions to a nucleic acid molecule encoding *Zea mays* AP1 (SEQ ID NO: 7), or a complementary sequence thereof.

In general, a nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule is a single-stranded nucleic acid sequence that can range in size from about 10 nucleotides to the full-length of a gene or a cDNA. Such a nucleotide sequence can be chemically synthesized, using routine methods or can be purchased from a commercial source. In addition, such nucleotide sequences can be obtained by enzymatic methods such as random priming methods, the polymerase chain reaction (PCR) or by standard restriction endonuclease digestion, followed by denaturation (Sambrook et al., *supra*, 1989).

A nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule can be used, for example, as a primer for PCR (Innis et al. (ed.) PCR Protocols: A Guide to Methods and Applications, San Diego, CA: Academic Press, Inc. 1990). Such a nucleotide sequence generally contains about 10 to about 50 nucleotides.

A nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule also can be used to screen a cDNA or genomic library to obtain a related nucleotide sequence. For example, a cDNA library that is prepared from rice or wheat can be screened with a nucleotide sequence derived from the *Zea mays* AP1 sequence in order to isolate a rice

or wheat ortholog of AP1. Generally, such a nucleotide sequence contains at least about 14-16 nucleotides depending, for example, on the hybridization conditions to be used.

5 A nucleotide sequence derived from a nucleic acid molecule encoding *Zea mays* AP1 (SEQ ID NO: 7) also can be used to screen a *Zea mays* cDNA library to isolate a sequence that is related to but distinct from AP1. Furthermore, such a hybridizing nucleotide sequence can
10 be used to analyze RNA levels or patterns of expression, as by northern blotting or by *in situ* hybridization to a tissue section. Such a nucleotide sequence also can be used in Southern blot analysis to evaluate gene structure and identify the presence of related gene sequences.

15 One skilled in the art would select a particular nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule encoding a floral meristem identity gene product based on the application for which the sequence will be
20 used. For example, in order to isolate an ortholog of AP1, one can choose a region of AP1 that is highly conserved among known AP1 sequences such as *Arabidopsis* AP1 (SEQ ID NO: 1) and *Zea mays* AP1 (GenBank accession number L46400; SEQ ID NO: 7). Similarly, in order to
25 isolate an ortholog of CAL, one can choose a region of CAL that is highly conserved among known CAL cDNAs, such as *Arabidopsis* CAL (SEQ ID NO: 9) and *Brassica* CAL (SEQ ID NO: 11). It further would be recognized, for example, that the region encoding the MADS domain, which is common

to a number of genes, can be excluded from the nucleotide sequence. In addition, one can use a full-length *Arabidopsis API* or *CAL* cDNA nucleotide sequence (SEQ ID NO: 1 or SEQ ID NO: 9) to isolate an ortholog of *API* or
5 *CAL*.

For example, the *Arabidopsis API* cDNA shown in Figure 1 (SEQ ID NO: 1) can be used as a probe to identify and isolate a novel *API* ortholog. Similarly, the *Arabidopsis CAL* cDNA shown in Figure 5 (SEQ ID NO: 9)
10 can be used to identify and isolate a novel *CAL* ortholog (see Examples IA and IIIC, respectively). In order to identify related MADS domain genes, a nucleotide sequence derived from the MADS domain of *API* or *CAL*, for example,
15 also can be useful to isolate a related gene sequence encoding this DNA-binding motif.

Hybridization utilizing a nucleotide sequence of the invention requires that hybridization be performed under relatively stringent conditions such that non-specific hybridization is minimized. Appropriate
20 hybridization conditions can be determined empirically, or can be estimated based, for example, on the relative G+C content of the probe and the number of mismatches between the probe and target sequence, if known. Hybridization conditions can be adjusted as desired by
25 varying, for example, the temperature of hybridizing or the salt concentration (Sambrook, *supra*, 1989).

The invention also provides a vector containing a nucleic acid molecule encoding a *CAL* gene product. In

addition, the invention provides a vector containing a nucleic acid molecule encoding the *Zea mays* AP1 gene product. A vector can be a cloning vector or an expression vector and provides a means to transfer an exogenous nucleic acid molecule into a host cell, which can be a prokaryotic or eukaryotic cell. Such vectors are well known and include plasmids, phage vectors and viral vectors. Various vectors and methods for introducing such vectors into a cell are described, for example, by Sambrook et al., *supra*, 1989, and by Glick and Thompson (eds.), Methods in Plant Molecular Biology and Biotechnology, Boca Raton, FL: CRC Press (1993), which is incorporated herein by reference.

The invention also provides an expression vector containing a nucleic acid molecule encoding a floral meristem identity gene product such as CAL, AP1 or LFY. Expression vectors are well known in the art and provide a means to transfer and express an exogenous nucleic acid molecule into a host cell. Thus, an expression vector contains, for example, transcription start and stop sites such as a TATA sequence and a poly-A signal sequence, as well as a translation start site such as a ribosome binding site and a stop codon, if not present in the coding sequence.

An expression vector can contain, for example, a constitutive regulatory element useful for promoting expression of an exogenous nucleic acid molecule in a plant cell. The use of a constitutive regulatory element can be particularly advantageous because expression from

the element is relatively independent of developmentally regulated or tissue-specific factors. For example, the cauliflower mosaic virus 35S promoter (CaMV35S) is a well-characterized constitutive regulatory element that

5 produces a high level of expression in all plant tissues (Odell et al., Nature 313:810-812 (1985), which is incorporated herein by reference). The CaMV35S promoter is particularly useful because it is active in numerous different angiosperms (Benfey and Chua, Science

10 250:959-966 (1990), which is incorporated herein by reference; Odell et al., *supra*, 1985). Other constitutive regulatory elements useful for expression in an angiosperm include, for example, the nopaline synthase (*nos*) gene promoter (An, Plant Physiol. 81:86 (1986),

15 which is herein incorporated by reference).

In addition, an expression vector of the invention can contain a regulated gene regulatory element such as a promoter or enhancer element. A particularly useful regulated promoter is a tissue-specific promoter

20 such as the shoot meristem-specific *CDC2* promoter (Hemerly et al., Plant Cell 5:1711-1723 (1993), which is incorporated herein by reference), or the *AGL8* promoter, which is active in the apical shoot meristem immediately after the transition to flowering (Mandel and Yanofsky,

25 Plant Cell 7:1763-1771 (1995), which is incorporated herein by reference).

An expression vector of the invention also can contain an inducible regulatory element, which has conditional activity dependent upon the presence of a

particular regulatory factor. Useful inducible regulatory elements include, for example, a heat-shock promoter (Ainley and Key, Plant Mol. Biol. 14:949 (1990), which is herein incorporated by reference) or a

5 nitrate-inducible promoter derived from the spinach nitrite reductase gene (Back et al., Plant Mol. Biol. 17:9 (1991), which is herein incorporated by reference). A hormone-inducible element

(Yamaguchi-Shinozaki et al., Plant Mol. Biol. 15:905

10 (1990) and Kares et al., Plant Mol. Biol. 15:225 (1990), which are herein incorporated by reference) or a light-inducible promoter, such as that associated with the small subunit of RuBP carboxylase or the LHCP gene families (Feinbaum et al., Mol. Gen. Genet. 226:449

15 (1991) and Lam and Chua, Science 248:471 (1990), which are herein incorporated by reference) also can be useful in an expression vector of the invention. A human glucocorticoid response element also can be used to achieve steroid hormone-dependent gene expression in

20 plants (Schena et al., Proc. Natl. Acad. Sci. USA 88:10421 (1991), which is herein incorporated by reference).

An appropriate gene regulatory element such as a promotor is selected depending on the desired pattern

25 or level of expression of a nucleic acid molecule linked thereto. For example, a constitutive promoter, which is active in all tissues, would be appropriate to express a desired gene product in all cells containing the vector. In addition, it can be desirable to restrict expression

30 of a nucleic acid molecule to a particular tissue or

during a particular stage of development. A developmentally regulated or tissue-specific expression can be useful for this purpose and can avoid potential undesirable side-effects that can accompany unregulated expression. Inducible expression also can be particularly useful to manipulate the timing of gene expression such that, for example, a population of transgenic angiosperms of the invention that contain an expression vector comprising a floral meristem identity gene linked to an inducible promoter can be induced to flower essentially at the same time. Such timing of flowering can be useful, for example, for manipulating the time of crop harvest.

The invention also provides a kit containing an expression vector having a nucleic acid molecule encoding a floral meristem identity gene product. Such a kit is useful for converting shoot meristem to floral meristem in an angiosperm or for promoting early flowering in an angiosperm. If desired, such a kit can contain appropriate reagents, which can allow relatively high efficiency of transformation of an angiosperm with the vector. Furthermore, a control plasmid lacking the floral meristem identity gene can be included in the kit to determine, for example, the efficiency of transformation.

The invention further provides a host cell containing a vector comprising a nucleic acid molecule encoding CAL. A host cell can be prokaryotic or eukaryotic and can be, for example, a bacterial cell,

yeast cell, insect cell, *xenopus* cell, mammalian cell or plant cell.

The invention also provides a transgenic garden variety cauliflower plant containing an exogenous nucleic acid molecule selected from the group consisting of a nucleic acid molecule encoding a CAL gene product and a nucleic acid molecule encoding an AP1 gene product. Such a transgenic cauliflower plant can produce an edible flower in place of the typical cauliflower vegetable.

A nucleic acid encoding CAL has been isolated from a *Brassica oleracea* line that produces wild-type flowers (*BoCAL*) and from the common garden variety of cauliflower, *Brassica oleracea* var. *botrytis* (*BobCAL*), which lacks flowers. The *Brassica oleracea* CAL cDNA (SEQ ID NO: 10) is highly similar to the *Arabidopsis* CAL cDNA (SEQ ID NO: 12; and see Figure 8). In contrast, the *Brassica oleracea* var. *botrytis* CAL cDNA contains a stop codon, predicting that the BobCAL protein will be truncated after amino acid 150 (SEQ ID NO: 14 and see Figure 8). The correlation of full-length *Arabidopsis* and *Brassica oleracea* CAL gene products with a flowering phenotype indicates that transformation of non-flowering garden varieties of cauliflower such as *Brassica oleracea* var. *botrytis* with a full-length CAL cDNA can induce flowering in the transgenic cauliflower plant.

As used herein, the term "CAL gene product" means a full-length CAL gene product that does not terminate substantially before amino acid 255 and that,

when ectopically expressed in shoot meristem, converts shoot meristem to floral meristem. A nucleic acid molecule encoding a CAULIFLOWER gene product can be, for example, a nucleic acid molecule encoding *Arabidopsis* CAL shown in Figure 5 (SEQ ID NO: 9) or a nucleic acid molecule encoding *Brassica oleracea* CAL shown in Figure 6 (SEQ ID NO: 11). In comparison, a nucleic acid molecule encoding a truncated CAL gene product that terminates substantially before amino acid 255, such as the encoded truncated BobCAL gene product (SEQ ID NO: 13), is not a nucleic acid molecule encoding a CAL gene product as defined herein. Furthermore, ectopic expression of BobCAL in an angiosperm does not result in conversion of shoot meristem to floral meristem.

As used herein, the term "AP1 gene product" means a full-length AP1 gene product that does not terminate substantially before amino acid 256. A nucleic acid molecule encoding an AP1 gene product can be, for example, a nucleic acid molecule encoding *Arabidopsis* AP1 shown in Figure 1 (SEQ ID NO: 1), *Brassica oleracea* AP1 shown in Figure 2, (SEQ ID NO: 3), *Brassica oleracea* var. *botrytis* AP1 shown in Figure 3 (SEQ ID NO: 5) or *Zea mays* AP1 shown in Figure 4 (SEQ ID NO: 7).

The invention provides a CAL polypeptide having at least about 70 percent amino acid identity with amino acids 1 to 160 of SEQ ID NO: 10 or SEQ ID NO: 12. For example, the *Arabidopsis thaliana* CAL polypeptide, having the amino acid sequence shown as amino acids 1 to 255 in Figure 5 (SEQ ID NO: 10), and the *Brassica oleracea* CAL

polypeptide, having the amino acid sequence shown as amino acids 1 to 255 in Figure 6 (SEQ ID NO: 12) are provided by the invention.

The invention also provides the truncated
5 *Brassica oleracea var. botrytis* CAL polypeptide having the amino acid sequence shown as amino acids 1 to 150 in Figure 7 (SEQ ID NO: 14). The BobCAL polypeptide can be useful as an immunogen to produce an antibody that specifically binds the truncated BoCAL polypeptide, but
10 does not bind a full length CAL gene product. Such an antibody can be useful to distinguish between a full length CAL and truncated CAL.

The invention provides also provides a *Zea mays* AP1 polypeptide. As used herein, the term "polypeptide" 15 is used in its broadest sense to include proteins, polypeptides and peptides, which are related in that each consists of a sequence of amino acids joined by peptide bonds. For convenience, the terms "polypeptide," "protein" and "gene product" are used interchangeably.
20 While no specific attempt is made to distinguish the size limitations of a protein and a peptide, one skilled in the art would understand that proteins generally consist of at least about 50 to 100 amino acids and that peptides generally consist of at least two amino acids up to a few
25 dozen amino acids. The term polypeptide is used generally herein to include any such amino acid sequence.

The term polypeptide also includes an active fragment of a floral meristem identity gene product. As

used herein, the term "active fragment," means a polypeptide portion of a floral meristem identity gene product that can convert shoot meristem to floral meristem or can provide early flowering. For example, an active fragment of a CAL polypeptide can consist of an amino acid sequence derived from a CAL protein as shown in Figure 5 or 6 (SEQ ID NOS: 10 and 12) and that has an activity of a CAL. An active fragment can be, for example, an amino terminal or carboxyl terminal truncated form of *Arabidopsis thaliana* CAL or *Brassica oleracea* CAL (SEQ ID NOS: 10 or 12, respectively). Such an active fragment can be produced using well known recombinant DNA methods (Sambrook et al., *supra*, 1989). The product of the *BobCAL* gene, which is truncated at amino acid 150, lacks activity in converting shoot meristem to floral meristem and, therefore, is an example of a polypeptide portion of a CAL floral meristem identity gene product that is not an "active fragment."

An active fragment of a floral meristem identity gene product can convert shoot meristem to floral meristem and is readily identified using the methods described in Example II, below). Briefly, *Arabidopsis* can be transformed with a nucleic acid molecule encoding a portion of a floral meristem identity gene product, in order to determine whether the fragment can convert shoot meristem to floral meristem or promote early flowering and, therefore, has an activity of a floral meristem identity gene product.

The invention further provides an antibody that specifically binds a CAL polypeptide, an antibody that specifically binds the truncated *Brassica oleracea* var. *botrytis* CAL polypeptide, and an antibody that

5 specifically binds the *Zea mays* API polypeptide. As used herein, the term "antibody" is used in its broadest sense to include polyclonal and monoclonal antibodies, as well as polypeptide fragments of antibodies that retain a specific binding activity for CAL protein of at least

10 about $1 \times 10^5 \text{ M}^{-1}$. One skilled in the art would know that anti-CAL antibody fragments such as Fab, F(ab'), and Fv fragments can retain specific binding activity for CAL and, thus, are included within the definition of an antibody. In addition, the term "antibody" as used

15 herein includes naturally occurring antibodies as well as non-naturally occurring antibodies and fragments that have binding activity such as chimeric antibodies or humanized antibodies. Such non-naturally occurring antibodies can be constructed using solid phase peptide

20 synthesis, produced recombinantly or obtained, for example, by screening combinatorial libraries consisting of variable heavy chains and variable light chains as described by Huse et al., *Science* 246:1275-1281 (1989), which is incorporated herein by reference.

25 An antibody "specific for" a polypeptide, or that "specifically binds" a polypeptide, binds with substantially higher affinity to that polypeptide than to an unrelated polypeptide. An antibody specific for a polypeptide also can have specificity for a related

30 polypeptide. For example, an antibody specific for

Arabidopsis CAL also can have specificity for *Brassica oleracea* CAL.

An anti-CAL antibody, for example, can be prepared using a CAL fusion protein or a synthetic peptide encoding a portion of *Arabidopsis* CAL or of *Brassica oleracea* CAL as an immunogen. One skilled in the art would know that purified CAL protein, which can be prepared from natural sources or produced recombinantly, or fragments of CAL, including a peptide portion of CAL such as a synthetic peptide, can be used as an immunogen. Non-immunogenic fragments or synthetic peptides of CAL can be made immunogenic by coupling the hapten to a carrier molecule such as bovine serum albumin (BSA) or keyhole limpet hemocyanin (KLH). In addition, various other carrier molecules and methods for coupling a hapten to a carrier molecule are well known in the art and described, for example, by Harlow and Lane, Antibodies: A laboratory manual (Cold Spring Harbor Laboratory Press, 1988), which is incorporated herein by reference. An antibody that specifically binds the truncated *Bob* CAL polypeptide or an antibody that specifically binds the *Zea mays* AP1 polypeptide similarly can be produced using such methods. An antibody that specifically binds the truncated *Brassica oleracea* var. *botrytis* CAL polypeptide can be particularly useful to distinguish between full-length CAL polypeptide and truncated CAL polypeptide.

The invention provides a method of identifying a *Brassica* having a modified *CAL* allele by detecting a polymorphism associated with a *CAL* locus, where the *CAL* locus comprises a modified *CAL* allele that does not encode an active *CAL* gene product. Such a method is useful for the genetic improvement of *Brassica* plants, a genus of great economic value.

Brassica plants are a highly diverse group of crop plants useful as vegetables and as sources of condiment mustard, edible and industrial oil, animal fodder and green manure. *Brassica* crops encompass a variety of well known vegetables including cabbage, cauliflower, broccoli, collard, kale, mustard greens, Chinese cabbage and turnip, which can be interbred for crop improvement (see, for example, King, Euphytica 50:97-112 (1990) and Crisp and Tapsell, Genetic improvement of vegetable crops pp. 157-178 (1993), each of which is herein incorporated by reference).

Breeding of *Brassica* crops is useful, for example, for improving the quality and early development of vegetables. In addition, such breeding can be useful to increase disease resistance, such as resistance, of a *Brassica* to clubroot disease or mildew; viral resistance, such as resistance to turnip mosaic virus and cauliflower mosaic virus; or pest resistance (King, *supra*, 1990).

The use of polymorphic molecular markers in the breeding of *Brassicae* is well recognized in the art (Crisp and Tapsell, *supra*, 1993). Identification of a

polymorphic molecular marker that is associated with a desirable trait can vastly accelerate the time required to breed the desirable trait into a new *Brassica* species or variant. In particular, since many rounds of 5 backcrossing are required to breed a new trait into a different genetic background, early detection of a desirable trait by molecular methods can be performed prior to the time a plant is fully mature, thus accelerating the rate of crop breeding (see, for example, 10 Figidore et al., *Euphytica* 69: 33-44 (1993), which is herein incorporated by reference).

A polymorphism associated with a *CAL* locus comprising a modified *CAL* allele that does not encode an active *CAL* gene product, is disclosed herein. Figure 6 15 shows the nucleotide (SEQ ID NO: 11) and amino acid (SEQ ID NO: 12) sequence of *Brassica oleracea* *CAL* (BoCAL), and Figure 7 shows the nucleotide (SEQ ID NO: 13) and amino acid (SEQ ID NO: 14) sequence of *Brassica oleracea* var. *botrytis* *CAL* (BobCAL). At amino acid 150, which is 20 glutamic acid (Glu) in BoCAL, a stop codon is present in BobCAL. This polymorphism results in a truncated BobCAL gene product that is not active as a floral meristem identity gene product. The BoCAL nucleic acid sequence (ACGAGT) can be readily distinguished from the BobCAL 25 nucleic acid sequence (ACTAGT) using well known molecular methods. For example, the polymorphic ACTAGT BobCAL sequence is recognized by a SpeI restriction endonuclease site, whereas the ACGAGT BoCAL sequence is not recognized by SpeI. Thus, a restriction fragment length 30 polymorphism (RFLP) in BobCAL provides a simple means for

identifying a modified *CAL* allele (*BobCAL*) and, therefore, can serve as a marker to predict the inheritance of the "cauliflower" phenotype.

A modified *CAL* allele encoding a truncated *CAL* gene product also can serve as a marker to predict the "cauliflower" phenotype in other cauliflower variants. For example, nine *romanesco* variants of *Brassica oleracea* var. *botrytis*, which each have the "cauliflower" phenotype, were examined for the presence of a stop codon at position 151 of the *CAL* coding sequence. All nine of the *romanesco* variants contained the *SpeI* site that indicates a stop codon and, thus, a truncated *CAL* gene product. In contrast, *Brassica oleracea* variants that lack the "cauliflower" phenotype (broccoli and brussels sprouts) were examined for the *SpeI* site. In every case, the broccoli and brussel sprout variants had a full-length *CAL* coding sequence, as indicated by the absence of the distinguishing *SpeI* site. Thus, a truncated *CAL* gene product can be involved in the "cauliflower phenotype" in numerous different *Brassica* variants.

As used herein, the term "modified *CAL* allele" means a *CAL* allele that does not encode a *CAL* gene product active in converting shoot meristem to floral meristem. A modified *CAL* allele can have a modification within a gene regulatory element such that a *CAL* gene product is not produced. In addition, a modified *CAL* allele can have a modification such as a mutation, deletion or insertion in a *CAL* coding sequence which

results in an inactive CAL gene product. For example, an inactive CAL gene product can result from a mutation creating a stop codon, such that a truncated, inactive CAL gene product lacking the ability to convert shoot
5 meristem to floral meristem is produced.

As used herein, the term "associated" means closely linked and describes the tendency of two genetic loci to be inherited together as a result of their proximity. If two genetic loci are associated and are
10 polymorphic, one locus can serve as a marker for the inheritance of the second locus. Thus, a polymorphism associated with a CAL locus comprising a modified CAL allele can serve as a marker for inheritance of the modified CAL allele. An associated polymorphism can be
15 located in proximity to a CAL gene or can be located within a CAL gene.

A polymorphism in a nucleic acid sequence can be detected by a variety of methods. For example, if the polymorphism occurs in a particular restriction
20 endonuclease site, the polymorphism can be detected by a difference in restriction fragment length observed following restriction with the particular restriction endonuclease and hybridization with a nucleotide sequence that is complementary to a nucleic acid sequence
25 including a polymorphism.

The use of restriction fragment length polymorphism as an aid to breeding *Brassicaceae* is well known in the art (see, for example, Slocum et al., Theor.

Appl. Genet. 80:57-64 (1990); Kennard et al., Theor. Appl. Genet. 87:721-732 (1994); and Figidore et al., *supra*, 1993, each of which is herein incorporated by reference). A restriction endonuclease such as SpeI, 5 which is useful for identifying the presence of a BobCAL allele in an angiosperm, is readily available and can be purchased from a commercial source. Furthermore, a nucleotide sequence that is complementary to a nucleic acid sequence having a polymorphism associated with a CAL 10 locus comprising a modified CAL allele can be derived, for example, from the nucleic acid molecule encoding *Brassica oleracea* var. *botrytis* CAL shown in Figure 7 (SEQ ID NO: 13) or from the nucleic acid molecule encoding *Brassica oleracea* CAL shown in Figure 6 (SEQ ID NO: 11).

In some cases, a polymorphism is not distinguishable by a RFLP, but nevertheless can be used to identify a *Brassica* having a modified CAL allele. For example, the polymerase chain reaction (PCR) can be used 20 to detect a polymorphism associated with a CAL locus comprising a modified CAL allele. Specifically, a polymorphic region of a modified allele can be selectively amplified by using a primer that matches the nucleotide sequence of one allele of a polymorphic locus, 25 but does not match the sequence of the second allele (Sobral and Honeycutt, The Polymerase Chain Reaction, pp. 304-319 (1994), which is herein incorporated by reference). Other well-known approaches for analyzing a polymorphism using PCR include discriminant hybridization 30 of PCR-amplified DNA to allele-specific oligonucleotides

and denaturing gradient gel electrophoresis (see Innis et al., *supra*, 1990).

The invention further provides a nucleic acid molecule encoding a chimeric protein, comprising a

5 nucleic acid molecule encoding a floral meristem identity gene product such as AP1, LFY or CAL operably linked to a nucleic acid molecule encoding a ligand binding domain. Expression of a chimeric protein of the invention in an angiosperm is particularly useful because the ligand

10 binding domain confers regulatable activity on a gene product such as a floral meristem identity gene product to which it is fused. Specifically, the floral meristem identity gene product component of the chimeric protein is inactive in the absence of the particular ligand,

15 whereas, in the presence of ligand, the ligand binds the ligand binding domain, resulting in floral meristem identity gene product activity.

A nucleic acid molecule encoding a chimeric protein of the invention contains a nucleic acid molecule

20 encoding a floral meristem identity gene product, such as a nucleic acid molecule encoding the amino acid sequence shown in Figure 1 (SEQ ID NO: 2), in Figure 5 (SEQ ID NO: 10), or in Figure 9 (SEQ ID NO: 10), either of which is operably linked to a nucleic acid molecule encoding a

25 ligand binding domain. The expression of such a nucleic acid molecule results in the production of a chimeric protein comprising a floral meristem identity gene product fused to a ligand binding domain. Thus, the invention also provides a chimeric protein comprising a

floral meristem identity gene product fused to a ligand binding domain.

A ligand binding domain useful in a chimeric protein of the invention can be a steroid binding domain such as the ligand binding domain of a glucocorticoid receptor, estrogen receptor, progesterone receptor, androgen receptor, thyroid receptor, vitamin D receptor or retinoic acid receptor. A particularly useful ligand binding domain is a glucocorticoid receptor ligand binding domain, encompassed, for example, within amino acids 512 to 795 of the rat glucocorticoid receptor as shown in Figure 16 (SEQ ID NO: 24; Miesfeld et al., Cell 46:389-399 (1986), which is incorporated herein by reference).

A chimeric protein containing a ligand binding domain, such as the rat glucocorticoid receptor ligand binding domain, confers glucocorticoid-dependent activity on the chimeric protein. For example, the activity of chimeric proteins consisting of adenovirus E1A, c-myc, c-fos, the HIV-1 Rev transactivator, MyoD or maize regulatory factor R fused to the rat glucocorticoid receptor ligand binding domain is regulated by glucocorticoid hormone (Eilers et al., Nature 340:66 (1989); Superti-Furga et al., Proc. Natl. Acad. Sci., U.S.A. 88:5114 (1991); Hope et al., Proc. Natl. Acad. Sci., U.S.A. 87:7787 (1990); Hollenberg et al., Proc. Natl. Acad. Sci., U.S.A. 90:8028 (1993), each of which is incorporated herein by reference).

Such a chimeric protein also can be regulated in plants. For example, a chimeric protein containing a heterologous protein fused to a rat glucocorticoid receptor ligand binding domain (amino acids 512 to 795) 5 was expressed under the control of the constitutive cauliflower mosaic virus 35S promoter in *Arabidopsis*. The activity of the chimeric protein was inducible; the chimeric protein was inactive in the absence of ligand, and became active upon treatment of transformed plants 10 with a synthetic glucocorticoid, dexamethasone (Lloyd et al., *Science* 266:436-439 (1994), which is incorporated herein by reference). As disclosed herein, a ligand binding domain fused to a floral meristem identity gene product can confer ligand inducibility on the activity of 15 a fused floral meristem identity gene product in plants such that, upon exposure to a particular ligand, the floral meristem identity gene product is active.

Methods for constructing a nucleic acid molecule encoding a chimeric protein are routine and well 20 known in the art (Sambrook et al., *supra*, 1989). For example, the skilled artisan would recognize that a stop codon in the 5' nucleic acid molecule must be removed and that the two nucleic acid molecules must be linked such that the reading frame of the 3' nucleic acid molecule is 25 preserved. Methods of transforming plants with nucleic acid molecules also are well known in the art (see, for example, Mohoney et al., U.S. patent number 5,463,174, and Barry et al., U.S. patent number 5,463,175, each of which is incorporated herein by reference).

As used herein, the term "operably linked," when used in reference to two nucleic acid molecules comprising a nucleic acid molecule encoding a chimeric protein, means that the two nucleic acid molecules are
5 linked in frame such that a full-length chimeric protein can be expressed. In particular, the 5' nucleic acid molecule, which encodes the amino-terminal portion of the chimeric protein, must be linked to the 3' nucleic acid molecule, which encodes the carboxyl-terminal portion of
10 the chimeric protein, such that the carboxyl-terminal portion of the chimeric protein is produced in the correct reading frame.

The invention further provides a transgenic angiosperm containing a nucleic acid molecule encoding a chimeric protein, comprising a nucleic acid molecule
15 encoding a floral meristem identity gene product such as AP1, CAL or LFY linked to a nucleic acid molecule encoding a ligand binding domain. Such a transgenic angiosperm is particularly useful because the angiosperm
20 can be induced to flower by contacting the angiosperm with a ligand that binds the ligand binding domain. Thus, the invention provides a method of promoting early flowering or of converting shoot meristem to floral meristem in a transgenic angiosperm containing a nucleic
25 acid molecule encoding a chimeric protein of the invention, comprising expressing the nucleic acid molecule encoding the chimeric protein in the angiosperm, and contacting the angiosperm with a ligand that binds the ligand binding domain, wherein binding of the ligand
30 to the ligand binding domain activates the floral

meristem identity gene product. In particular, the invention provides methods of promoting early flowering or of converting shoot meristem to floral meristem in a transgenic angiosperm containing a nucleic acid molecule 5 encoding a chimeric protein that consists of a nucleic acid molecule encoding AP1 or CAL or LFY linked to a nucleic acid molecule encoding a glucocorticoid receptor ligand binding domain by contacting the transgenic angiosperm with a glucocorticoid such as dexamethasone.

10 As used herein, the term "ligand" means a naturally occurring or synthetic chemical or biological molecule such as a simple or complex organic molecule, a peptide, a protein or an oligonucleotide that specifically binds a ligand binding domain. A ligand of 15 the invention can be used, alone, in solution or can be used in conjunction with an acceptable carrier that can serve to stabilize the ligand or promote absorption of the ligand by an angiosperm.

One skilled in the art can readily determine 20 the optimum concentration of ligand needed to bind a ligand binding domain and render a floral meristem identity gene product active. Generally, a concentration of about 1 nM to 1 μ M dexamethasone is useful for activating floral meristem identity gene product activity 25 in a chimeric protein comprising a floral meristem identity gene product and a glucocorticoid receptor ligand binding domain (Lloyd et al., *supra*, 1994).

A transgenic angiosperm expressing a chimeric protein of the invention can be contacted with ligand in a variety of manners including, for example, by spraying, injecting or immersing the angiosperm. Further, a plant 5 may be contacted with a ligand by adding the ligand to the plant's water supply or to the soil, whereby the ligand is absorbed into the angiosperm.

The following examples are intended to 10 illustrate but not limit the present invention.

EXAMPLE I

Identification and characterization of the Zea mays APETALA1 cDNA

This example describes the isolation and 15 characterization of the *Zea mays* ZAP-1 "gene", which is an ortholog of the *Arabidopsis* floral meristem identity gene, *AP1*.

A. Identification and characterization of a nucleic acid sequence encoding ZAP-1

20 The utility of using a cloned floral homeotic gene from *Arabidopsis* to identify the putative ortholog in maize has previously been demonstrated (Schmidt et al., *supra*, (1993), which is incorporated herein by reference). As described in Mena et al. (*Plant J.* 25 8(6):845-854 (1995)), the maize ortholog of the *Arabidopsis* *AP1* floral meristem identity gene, was isolated by screening a *Zea mays* ear cDNA library using

the *Arabidopsis API* cDNA (SEQ ID NO: 1) as a probe. A cDNA library was prepared from wild-type immature ears as described by Schmidt et al., *supra*, 1993, using an *Arabidopsis API* cDNA sequence as a probe. The 5 *Arabidopsis API* cDNA (SEQ ID NO: 1), which is shown in Figure 1 (SEQ ID NO 1), was used as the probe. Low-stringency hybridizations with the API probe were conducted as described previously for the isolation of ZAG1 using the AG cDNA as a probe (Schmidt et al., *supra*, 10 1993). Positive plaques were isolated and cDNAs were recovered in Bluescript by *in vivo* excision. Double-stranded sequencing was performed using the Sequenase Version 2.0 kit (U.S. Biochemical, Cleveland, Ohio) according to the manufacturer's protocol.

15 The cDNA sequence and deduced amino acid sequence for ZAPI are shown in Figure 4 (SEQ ID NOS: 7 and 8). The deduced amino acid sequence for ZAPI shares 89% identity with *Arabidopsis API* through the MADS domain (amino acids 1 to 57) and 70% identity through the first 20 160 amino acids, which includes the K domain. The high level of amino acid sequence identity between ZAPI and API (SEQ ID NOS: 8 and 2), as well as the expression pattern of ZAPI in maize florets (see below), indicates that ZAPI is the maize ortholog of *Arabidopsis API*.

25 B. RNA expression pattern of ZAPI

Total RNA was isolated from different maize tissues as described by Cone et al., Proc. Natl. Acad. Sci. USA 83:9631-9635 (1986), which is herein

incorporated by reference. RNA was prepared from ears or tassels at early developing stages (approximately 2 cm in size), husk leaves from developing ear shoots, shoots and roots of germinated seedlings, leaves from 2 to 3 week old plants and endosperm, and embryos at 18 days after pollination. Mature floral organs were dissected from ears at the time of silk emergence or from tassels at several days pre-emergence. To study expression patterns in the mature female flower, carpels were isolated and the remaining sterile organs were pooled and analyzed together. In the same way, stamens were dissected and collected from male florets and the remaining organs (excluding the glumes) were pooled as one sample.

RNA concentration and purity was determined by absorbance at 260/280 nM, and equal amounts (10 µg) were fractionated on formaldehyde-agarose gels. Gels were stained in a solution of 0.125 µg ml⁻¹ acridine orange to confirm the integrity of the RNA samples and the uniformity of gel loading, then RNA was blotted on to Hybond-N® membranes (Amersham International, Arlington Heights, Illinois) according to the manufacturer's instructions. Prehybridization and hybridization solutions were prepared as previously described (Schmidt et al., Science 238:960-963 (1987), which is incorporated herein by reference). The probe for ZAPI RNA expression studies was a 445 bp SacI-NsiI fragment from the 3' end of the cDNA. Southern blot analyses were conducted to establish conditions for specific hybridization of this probe. No cross-hybridization was detected with

hybridization at 60°C in 50% formamide and washes at 65°C in 0.1x SSC and 0.5% SDS.

The strong sequence similarity between ZAP1 and API indicated that ZAP1 was the ortholog of this 5 *Arabidopsis* floral meristem identity gene. As a first approximation of whether the pattern of ZAP1 expression paralleled that of API, a blot of total RNA from vegetative and reproductive organs was hybridized with a gene-specific fragment of the ZAP1 cDNA (nucleotides 370 10 to 820 of SEQ ID NO: 7). ZAP1 RNA was detected only in male and female inflorescences and in the husk leaves that surround the developing ear. No ZAP1 RNA expression was detectable in RNA isolated from root, shoot, leaf, endosperm, or embryo tissue. The restriction of ZAP1 15 expression to terminal and axillary inflorescences is consistent with ZAP1 being the *Arabidopsis* API ortholog.

Male and female florets were isolated from mature inflorescences, and the reproductive organs were separated from the remainder of the floret. RNA was 20 isolated from the reproductive and the sterile portions of the florets. ZAP1 RNA expression was not detected in maize stamens or carpels, whereas high levels of ZAP1 RNA were present in developing ear and tassel florets from which the stamens and carpels had been removed. 25 Thus, the exclusion of ZAP1 expression in stamens and carpels and its inclusion in the RNA of the non-reproductive portions of the floret (lodicles, lemma and palea) is similar to the pattern of expression of API in flowers of *Arabidopsis*.

EXAMPLE II**Conversion of shoot meristem to floral meristem in an APETALA1 transgenic plant**

This example describes methods for producing a
5 transgenic *Arabidopsis* plant, in which shoot meristem is
converted to floral meristem.

A. Ectopic expression of APETALA1 converts inflorescence shoots into flowers

Transgenic plants that constitutively express
10 *API* from the cauliflower mosaic virus 35S (CaMV35S)
promoter were produced to determine whether ectopic *API*
expression could convert shoot meristem to floral
meristem. The *API* coding sequence was placed under
control of the cauliflower mosaic virus 35S promoter
15 (Odell et al., *supra*, 1985) as follows. BamHI linkers
were ligated to the HincII site of the full-length *API*
complementary DNA (Mandel et al., *supra*, (1992), which is
incorporated herein by reference) in pAM116, and the
resulting BamHI fragment was fused to the cauliflower
20 mosaic virus 35S promoter (Jack et al., *Cell* 76:703-716
(1994), which is incorporated herein by reference) in
pCGN18 to create pAM563.

Transgenic *API* *Arabidopsis* plants of the
Columbia ecotype were generated by selecting
25 kanamycin-resistant plants after *Agrobacterium*-mediated
plant transformation using the *in planta* method (Bechtold

et al., C.R. Acad. Sci. Paris 316:1194-1199 (1993), which is incorporated herein by reference). All analyses were performed in subsequent generations. Approximately 120 independent transgenic lines that displayed the described 5 phenotypes were obtained.

Remarkably, in 35S-AP1 transgenic plants, the normally indeterminate shoot apex) prematurely terminated as a floral meristem and formed a terminal flower. In addition, all lateral meristems that normally 10 would produce inflorescence shoots also were converted into solitary flowers. These results demonstrate that ectopic expression of AP1 in shoot meristem is sufficient to convert shoot meristem to floral meristem, even though AP1 normally is not absolutely required to specify floral 15 meristem identity.

B. LEAFY is not required for the conversion of inflorescence shoots to flowers in an APETALA1 transgenic plant

To determine whether the 35S-AP1 transgene 20 causes ectopic LFY activity, and whether ectopic LFY activity is required for the conversion of shoot meristem to floral meristem, the 35S-AP1 transgene was introduced into *Arabidopsis lfy* mutants. The 35S-AP1 transgene was crossed into the strong *lfy-6* mutant background and the F₂, 25 progeny were analyzed.

Lfy mutant plants containing the 35S-*API* transgene displayed the same conversion of apical and lateral shoot meristem to floral meristem as was observed in transgenics containing wild type *LFY*. However, the 5 resulting flowers had the typical *lfy* mutant phenotype, in which floral organs developed as sepaloid and carpelloid structures, with an absence of petals and stamens. These results demonstrate that *LFY* is not required for the conversion of shoot meristem to floral 10 meristem in a transgenic angiosperm that ectopically expresses *API*.

C. *APETALA1* is not sufficient to specify organ fate

As well as being involved in the early step of specifying floral meristem identity, *API* also is involved 15 in specifying sepal and petal identity at a later stage in flower development. Although *API* RNA is initially expressed throughout the young flower primordium, it is later excluded from stamen and carpel primordia (Mandel et al., *Nature* 360:273-277 (1992)). Since the 20 cauliflower mosaic virus 35S promoter is active in all floral organs, 35S-*API* transgenic plants are likely to ectopically express *API* in stamens and carpels. However, 35S-*API* transgenic plants had normal stamens and carpels, indicating that *API* is not sufficient to specify sepal 25 and petal organ fate.

D. Ectopic expression of APETALA1 causes early flowering

In addition to its ability to alter inflorescence meristem identity, ectopic expression of API also influences the vegetative phase of plant growth.

5 Wild-type plants have a vegetative phase during which a basal rosette of leaves is produced, followed by the transition to reproductive growth. The transition from vegetative to reproductive growth was measured both in terms of the number of days post-germination until the

10 first visible flowers were observed, and by counting the number of leaves. Under continuous light, wild-type and 35S-API transgenic plants flowered after producing 9.88 ± 1.45 and 4.16 ± 0.97 leaves, respectively. Under short-day growth conditions (8 hours light, 16 hours dark, 24 C),

15 wild-type and 35S-API transgenic plants flowered after producing 52.42 ± 3.47 and 7.4 ± 1.18 leaves, respectively.

In summary, under continuous light growth conditions, flowers appear on wild-type *Arabidopsis* plants after approximately 18 days, whereas the 35S-API transgenic plants flowered after an average of only 10 days. Furthermore, under short-day growth conditions, flowering is delayed in wild-type plants until approximately 10 weeks after germination, whereas, 35S-API transgenic plants flowered in less than 3 weeks.

25 Thus, ectopic API activity significantly reduced the time to flowering and reduced the delay of flowering caused by short day growth conditions.

EXAMPLE III**Isolation and characterization of the *Arabidopsis* and *Brassica oleracea* CAULIFLOWER genes**

This example describes methods for isolating
5 and characterizing the *Arabidopsis* and *Brassica oleracea*
CAL genes.

A. Isolation of the *Arabidopsis* and *Brassica oleracea* CAULIFLOWER genes

Genetic evidence that *CAL* and *AP1* proteins may
10 be functionally related indicated that these proteins may
share similar DNA sequences. In addition, DNA blot
hybridization revealed that the *Arabidopsis* genome
contains a gene that is closely related to *AP1*. The *CAL*
gene, which is closely related to *AP1*, was isolated and
15 identified as a member of the family of *Arabidopsis* MADS
domain genes known as the *AGAMOUS-like* (*AGL*) genes.

Hybridization with an *AP1* probe was used to
isolate a 4.8-kb Eco RI genomic fragment of *CAL*. The
corresponding *CAL* complementary DNA (pBS85) was cloned by
20 reverse transcription-polymerase chain reaction (RT-PCR)
with the oligonucleotides AGL10-1
(5'-GATCGTCGTTATCTCTCTTG-3'; SEQ ID NO: 25) and AGL10-12
(5'-GTAGTCTATTCAAGCGGCG-3'; SEQ ID NO: 26).

The *Arabidopsis* *CAL* cDNA encodes a putative 255
25 amino acid protein (Figure 5; SEQ ID NO: 10) having a
calculated molecular weight of 30.1 kD and an isoelectric

point of 8.78. The deduced amino acid sequence for CAL contains a MADS domain which generally is present in a class of transcription factors. The MADS domains of CAL and AP1 were markedly similar, differing in only 5 of 56 5 amino acid residues, 4 of which represent conservative replacements. Overall, the putative CAL protein is 76% identical to AP1; with allowance for conservative amino acid substitutions, the two proteins are 88% similar. These results indicate that CAL and AP1 may recognize 10 similar target sequences and regulate many of the same genes involved in floral meristems identity.

CAL was mapped to the approximate location of the loci identified by classical genetic means for the cauliflower phenotype (Bowman et al., Development 119:721 15 (1993), which is herein incorporated by reference). Restriction fragment length polymorphism (RFLP) mapping filters were scored and the results analyzed with the Macintosh version of the Mapmaker program as described by Rieter et al., (Proc. Natl. Acad. Sci. USA, 89:1477 20 (1992), which is herein incorporated by reference). The results localized CAL to the upper arm of chromosome 1, near marker λ235.

A genomic fragment spanning the CAL gene was used to transform cal-1 ap1-1 plants. A 5850-bp Bam HI 25 fragment containing the entire coding region of the *Arabidopsis* CAL gene as well as 1860 bp upstream of the putative translational start site was inserted into the pBIN19 plant transformation vector (Clontech, Palo Alto, California) and used for transformation of root tissue

from *cal-1* *apl-1* plants as described by Valvekens et al. (Proc. Natl. Acad. Sci., USA 85:5536 (1988), which is incorporated herein by reference). Seeds were harvested from primary transformants, and all phenotypic analyses 5 were performed in subsequent generations. Four independent lines transformed with *CAL* showed a complementation of the cauliflower (*cal*) phenotype and displayed a range of phenotypes similar to those exhibited by *apl* mutants. These results demonstrated 10 that *CAL* functions to convert shoot meristem to floral meristem.

In order to identify regions of functional importance in the *CAL* protein, *cal* mutants were generated and analyzed. The *cal* alleles were isolated by 15 mutagenizing seeds homozygous for the *apl-1* allele in Ler with 0.1% or 0.05% ethylmethane sulfonate (EMS) for 16 hours. Putative new *cal* alleles were crossed to *cal-1* *apl-1* *chlorina* plants to verify allelism. Two sets of oligonucleotides were used to amplify and clone new 20 alleles: oligos AGL10-1 (SEQ ID NO: 25) and AGL10-2 (5'-GATGGAGACCATTAAACAT-3'; SEQ ID NO: 27) for the 5' portion and oligos AGL10-3 (5'-GGAGAAGGTACTAGAACG-3'; SEQ ID NO: 28) and AGL10-4 (5'-GCCCTCTTCCATAGATCC-3'; SEQ ID NO: 29) for the 3' portion of the gene. All coding 25 regions and intron-exon boundaries of the mutant alleles were sequenced.

Sequence analysis of the *cal-1* allele, which exists in the wild-type *Wassilewskija* (WS) ectotype, revealed a cluster of three amino acid differences in the

seventh exon, relative to the wild-type gene product from Landsberg erecta (Ler) (Figure 8). One or more of these amino acid differences can be responsible for the *cal* phenotype, because the *cal-1* gene was expressed normally 5 and the transcribed RNA was correctly spliced in the WS background. The three additional *cal* alleles that were isolated, designated *cal-2*, *cal-3*, and *cal-4*, exhibited phenotypes similar to that of the *cal-1* allele.

Sequence analyses revealed a single missense 10 mutation for each (Figure 8). Since mutations in the *cal-2* and *cal-3* alleles lie in the MADS domain, these mutations can affect the ability of CAL to bind DNA and activate its target genes. Because the *cal-4* allele contains a substitution in the K domain, a motif thought 15 to be involved in protein-protein interactions, this mutation can affect the ability of CAL to form homodimers or to interact with other proteins such as AP1.

B. RNA expression pattern of CAULIFLOWER

To characterize the temporal and spatial 20 pattern of *CAL* RNA accumulation, RNA *in situ* hybridizations were performed using a *CAL*-specific probe. ³⁵S-labeled antisense *CAL* and *BoCAL* mRNA was synthesized 25 from Sca 1-digested cDNA templates and hybridized to 8 µm sections of *Arabidopsis Ler* or *Brassica oleracea* inflorescences. The probes did not contain any MADS box sequences in order to avoid cross-hybridization with other MADS box genes. Hybridization conditions were as

previously described (Drews et al., *Cell* 65:991 (1991), which is herein incorporated by reference).

As with *API*, *CAL* RNA accumulated in young flower primordia, consistent with the ability of *CAL* to substitute for *API* in specifying floral meristems. In contrast to *API* RNA, however, which accumulated at high levels throughout sepal and petal development, *CAL* RNA was detected only at very low levels in these organs. These results demonstrate that *CAL* was unable to substitute for *API* in specifying sepals and petals, at least in part as a result of the relatively low levels of *CAL* RNA in these developing organs.

C. Molecular Basis of the cauliflower phenotype

The *cal* phenotype in *Arabidopsis* is similar to the inflorescence structure that develops in the closely related species *Brassica oleracea* var. *botrytis*, the cultivated garden variety of cauliflower, indicating that the *CAL* gene can contribute to the *cal* phenotype of this agriculturally important species. Thus, *CAL* gene homologs were isolated from a *Brassica oleracea* line that produces wild-type flowers (*BoCAL*) and from the common garden variety of cauliflower *Brassica oleracea* var. *botrytis* (*BobCAL*).

The single-copy *BobCAL* gene (Snowball Y Improved, NK Lawn & Garden, Minneapolis, MN) was isolated from a size-selected genomic library in λ BlueStar (Novagen) on a 16-kbp BamHI fragment with the *Arabidopsis*

CAL gene as a probe. The BoCAL gene was isolated from a rapid cycling line (Williams and Hill, *Science* 232:1385 (1986)) by PCR on both RNA and genomic DNA. The cDNA was isolated by RT-PCR using the oligonucleotides: Bob1
5 (5'-TCTACGAGAAATGGGAAGG-3'; SEQ ID NO: 30) and Bob2 (5'-GTCGATATATGGCGAGTCC-3'; SEQ ID NO: 31). The 5' portion of the gene was obtained using oligonucleotides Bob 1 (SEQ ID NO: 30) and Bob4B (5'-CCATTGACCAGTTCGTTG-3'; SEQ ID NO: 32). The 3'
10 portion was obtained using oligonucleotides Bob3 (5'-GCTCCAGACTCTCACGTC-3'; SEQ ID NO: 33) and Bob2 (SEQ ID NO: 31).

RNA *in situ* hybridizations were performed to determine the expression pattern of BoCAL gene from 15 *Brassica oleracea*. As in *Arabidopsis*, BoCAL RNA accumulated uniformly in early floral primordia and later was excluded from the cells that give rise to stamens and carpels.

DNA sequence analyses revealed that the open 20 reading frame of the BoCAL gene is intact, whereas that of the BobCAL gene is interrupted by a stop codon in exon 5 (Figure 8). Translation of the resulting BobCAL protein product is truncated after only 150 of the wild-type 255 amino acids. Because similar stop codon 25 mutations in the fifth exon of the *Arabidopsis* API coding sequence result in plants having a severe *api* phenotype, the BobCAL protein likely is not functional. These results indicate that, as in *Arabidopsis*, the molecular basis for the cauliflower phenotype in *Brassica oleracea*

var. *botrytis* is due, at least in part, to a mutation in the *BobCAL* gene.

EXAMPLE IV

5 Conversion of inflorescence shoots into flowers in an
CAULIFLOWER transgenic plant

This example describes methods for producing a transgenic CAL plant.

A. Ectopic expression of CAULIFLOWER converts
inflorescence shoots to flowers

10 Transgenic *Arabidopsis* plants that ectopically express *CAL* in shoot meristem were generated. The full-length *CAL* cDNA was inserted downstream of the 35S cauliflower mosaic virus promoter in the EcoRI of pMON530 (Monsanto Co. Co., St. Louis, Missouri). This plasmid was 15 introduced into *Agrobacterium* strain ASE (check) and used to transform the Columbia ecotype of *Arabidopsis* using a modified vacuum infiltration method described by Bechtold et al. (*supra*, 1993). The 96 lines generated that harbored the 35S-CAL construct had a range of weak to 20 strong phenotypes. The transgenic plants with the strongest phenotypes (27 lines) closely resembled the *tfl* mutant.

35S-CAL transgenic plants had converted apical and lateral inflorescence shoots into flowers and showed 25 an early flowering phenotype. These results demonstrate

that *CAL* is sufficient for the conversion of shoots to flowers and for promoting early flowering.

EXAMPLE V

5 Conversion of shoots into flowers in a
LEAFY transgenic plant

This example describes methods for producing a transgenic *LFY* *Arabidopsis* and aspen.

A. Conversion of Arabidopsis shoots by LEAFY

Transgenic *Arabidopsis* plants were generated by
10 transforming *Arabidopsis* with *LFY* under the control of
the cauliflower mosaic virus 35S promoter (CaMV35S) (Odell
et al., *supra*, (1985)). A *LFY* complementary cDNA (Weigel
et al., *Cell* 69:843-859 (1992), which is incorporated
herein by reference) was inserted into a T-DNA
15 transformation vector containing a CaMV 35S promoter/3'
nos cassette (Jack et al., *supra*, 1994). Transformed
seedlings were selected for kanamycin resistance.
Several hundred transformants in three different genetic
backgrounds (Nossen, Wassilewskija and Columbia) were
20 recovered and several lines were characterized in detail.

High levels of *LFY* RNA expression were detected
by northern blot analysis. In general, Nossen lines had
weaker phenotypes, especially when grown in short days.
The 35S-*LFY* transgene of line DW151.117 (ecotype
25 Wassilewskija) was introgressed into the *erecta*
background by backcrossing to a Landsberg *erecta* strain.

Plants were grown under 16 hours light and 8 hours dark. The *35S-LFY* transgene provided at least as much *LFY* activity as the endogenous gene and completely suppressed the *lfy* mutant phenotype when crossed into the background 5 of the *lfy-6* null allele.

Most *35S-LFY* transgenic plants lines demonstrated a very similar, dominant and heritable phenotype. Secondary shoots that arose in lateral positions were consistently replaced by solitary flowers, 10 and higher-order shoots were absent. Although the number of rosette leaves was unchanged from the wild type, *35S-LFY* plants flowered earlier than wild type; the solitary flowers in the axils of the rosette leaves developed and opened precociously. In addition, the 15 primary shoot terminated with a flower. In the most extreme cases, a terminal flower was formed immediately above the rosette. This gain of function phenotype (conversion of shoots to flowers) is the opposite of the *lfy* loss of function phenotype (conversion of flowers to 20 shoots). These results demonstrate that *LFY* encodes a developmental switch that is both sufficient and necessary to convert shoot meristem to flower meristem.

The effects of constitutive *LFY* expression differ for primary and secondary shoot meristems. 25 Secondary meristems were transformed into flower meristem, apparently as soon as it developed, and produced only a single, solitary flower. In contrast, primary shoot meristem produced leaves and lateral flowers before being consumed in the formation of a

terminal flower. These developmental differences indicate that a meristem must acquire competence to respond to the activity of a floral meristem identity gene such as *LFY*.

5 B. Conversion of aspen shoots by *LEAFY*

Given that constitutive expression of *LFY* induced precocious flowering during the vegetative phase of *Arabidopsis*, the effect of *LFY* on the flowering of other species was examined. The perennial tree, hybrid 10 aspen, is derived from parental species that flower naturally only after 8-20 years of growth (Schopmeyer (ed.), USDA Agriculture Handbook 450: Seeds of Woody Plants in the United States, Washington DC, USA: US Government Printing Office, pp. 645-655 (1974)). 35S-*LFY* 15 aspen plants were obtained by *Agrobacterium*-mediated transformation of stem segments and subsequent regeneration of transgenic shoots in tissue culture.

Hybrid aspen was transformed exactly as described by Nilsson et al. (Transgen. Res. 1:209-220 20 (1992), which is incorporated herein by reference). Levels of *LFY* RNA expression were similar to those of 35S-*LFY* *Arabidopsis*, as determined by northern blot analysis. The number of vegetative leaves varied between different regenerating shoots, and those with a higher 25 number of vegetative leaves formed roots, allowing for transfer to the greenhouse. Individual flowers were removed either from primary transformants that had been transferred to the greenhouse, or from catkins collected

in spring, 1995, at Carlshem, Umeå, Sweden) from a tree whose age was determined by counting the number of annual rings in a core extracted with an increment borer at 1.5 meters above ground level. Flowers were fixed in 5 formaldehyde/acetic acid/ethanol and destained in ethanol before photography.

The overall phenotype of *35S-LFY* aspen was similar to that of *35S-LFY* *Arabidopsis*. In wild-type plants of both species, flowers normally are formed in 10 lateral positions on inflorescence shoots. In aspen, these inflorescence shoots, called catkins, arise from the leaf axils of adult trees. In both *35S-LFY* *Arabidopsis* and *35S-LFY* aspen, solitary flowers were formed instead of shoots in the axils of vegetative 15 leaves. Moreover, as in *Arabidopsis*, the secondary shoots of transgenic aspen were more severely affected than the primary shoot.

Regenerating *35S-LFY* aspen shoots initially produced solitary flowers in the axils of normal leaves. 20 However, the number of vegetative leaves was limited, and the shoot meristem was prematurely consumed in the formation of an aberrant terminal flower. Precocious flower development was specific to *35S-LFY* transformants and was not observed in non-transgenic controls. 25 Furthermore, not a single instance of precocious flower development has been observed in more than 1,500 other lines of transgenic aspen generated with various constructs from 1989 to 1995 at the Swedish University of Agricultural Sciences. These results demonstrate that a

heterologous floral meristem identity gene product is active in an angiosperm.

Although the invention has been described with reference to the examples above, it should be understood 5 that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims.

We claim:

1. A nucleic acid molecule encoding a CAULIFLOWER (CAL) gene product having at least about 70 percent amino acid identity with amino acids 1 to 160 of 5 the sequence shown in Figure 5 (SEQ ID NO: 10) or with amino acids 1 to 160 of the sequence shown in Figure 6 (SEQ ID NO: 12).
2. The nucleic acid molecule of claim 1, wherein said CAL gene product is selected from the group 10 consisting of *Arabidopsis thaliana* CAL having the amino acid sequence shown in Figure 5 (SEQ ID NO: 10) and *Brassica oleracea* CAL having the amino acid sequence shown in Figure 6 (SEQ ID NO: 12).
3. A nucleic acid molecule selected from the 15 group consisting of a nucleic acid molecule having the nucleic acid sequence shown in Figure 5 (SEQ ID NO: 9) and a nucleic acid molecule having the nucleic acid sequence shown in Figure 6 (SEQ ID NO: 11).
4. A nucleic acid molecule encoding a 20 truncated CAL gene product having at least about 70 percent amino acid identity with amino acids 1 to 150 of the sequence shown in Figure 7 (SEQ ID NO: 14).
5. The nucleic acid molecule of claim 4, wherein said truncated CAL gene product is *Brassica oleracea* var. *botrytis* CAL having the amino acid sequence 25 shown in Figure 7 (SEQ ID NO: 14).
6. A nucleic acid molecule having the nucleic acid sequence shown in Figure 7 (SEQ ID NO: 13).

7. A nucleotide sequence that hybridizes under relatively stringent conditions to a nucleic acid molecule selected from the group consisting of:

5 the nucleic acid molecule of claim 3 or a nucleic acid molecule complementary thereto; and
 the nucleic acid molecule of claim 6 or a nucleic acid molecule complementary thereto.

10 8. A *CAL* gene, comprising a *CAL* gene selected from the group consisting of an *Arabidopsis thaliana* *CAL* gene having the nucleotide sequence shown in Figure 13 (SEQ ID NO: 20), a *Brassica oleracea* *CAL* gene having the nucleotide sequence shown in Figure 14 (SEQ ID NO: 21) 15 and a *Brassica oleracea* var. *botrytis* *CAL* gene having the nucleotide sequence shown in Figure 15 (SEQ ID NO: 22).

9. A nucleotide sequence that hybridizes under relatively stringent conditions to the *CAL* gene of claim 8, or a complementary sequence thereto.

20 10. A vector, comprising the nucleic acid molecule of claim 1.

11. A vector, comprising the gene of claim 8.

25 12. A vector, comprising a nucleic acid molecule selected from the group consisting of the nucleic acid molecule of claim 2 and the nucleic acid molecule of claim 3.

13. A host cell, comprising the vector of claim 10.

14. The vector of claim 10, wherein said vector is an expression vector.

15. An expression vector, comprising a nucleic acid molecule selected from the group consisting of the 5 nucleic acid molecule of claim 2 and the nucleic acid molecule of claim 3.

16. The expression vector of claim 14, further comprising a cauliflower mosaic virus 35S promoter.

17. The expression vector of claim 14, further 10 comprising an inducible regulatory element.

18. A kit for converting shoot meristem to floral meristem in an angiosperm, comprising the expression vector of claim 14.

19. A kit for promoting early flowering in an 15 angiosperm, comprising the expression vector of claim 14.

20. A CAL polypeptide having at least about 70 percent amino acid identity with amino acids 1 to 160 of the sequence shown in Figure 5 (SEQ ID NO: 10) or with amino acids 1 to 160 of the sequence shown in Figure 6 20 (SEQ ID NO: 12).

21. The CAL polypeptide of claim 20, wherein said CAL polypeptide is *Arabidopsis thaliana* CAL polypeptide having the amino acid sequence shown as amino acids 1 to 255 in Figure 5 (SEQ ID NO: 10).

22. The CAL polypeptide of claim 20, wherein said CAL polypeptide is *Brassica oleracea* CAL polypeptide having the amino acid sequence shown as amino acids 1 to 255 in Figure 6 (SEQ ID NO: 12).

5 23. An antibody that specifically binds the CAL polypeptide of claim 20.

24. The antibody of claim 23, wherein said antibody is a monoclonal antibody.

25. A truncated *Brassica oleracea* var. 10 *botrytis* CAL polypeptide having the amino acid sequence shown as amino acids 1 to 150 in Figure 7 (SEQ ID NO: 14).

26. An antibody that specifically binds the truncated *Brassica oleracea* var. *botrytis* CAL polypeptide 15 of claim 25.

27. A method of identifying a *Brassica* having a modified CAL allele, comprising detecting a polymorphism associated with a CAL locus, said CAL locus comprising a modified CAL allele that does not encode an 20 active CAL gene product.

28. The method of claim 27, wherein said modified CAL allele encodes a truncated CAL gene product.

29. The method of claim 27, wherein said polymorphism is within a CAL gene.

30. The method of claim 29, wherein said polymorphism is detectable as a restriction fragment length polymorphism.

31. The method of claim 30, wherein said polymorphism is at nucleotide 451 of the nucleic acid sequence shown in Figure 7 (SEQ ID NO: 13).

1 / 44

-81

GAATCCCTCG AGCTACGGTCA GGGCCCTGAC GTAGCTCGAA CTCTGAGCTC TCTTTATAT

-21

CTCTCTTGTA GTTCTTATT GGGGGCTCTT GTTTTGTGTTG GTTCCTTTAG AGTAAGAAGT

TCTTAAAAAA AGGATCAAAA ATG GCA AGG GGT AGG GTT CAA TTG AAG AGG ATA
M G R G R V Q L K R I> 11

40

GAG AAC AAG ATC AAT AGA CAA GTG ACA TTC TCG AAA AGA AGA GCT GGT
E N K I N R Q V T F S K R R A Q> 27

100

CTT TIG AAG AAA GCT CAT GAG ATC TCT GTT CTC TGT GAT GCT GAA GIT
L L K K A H E I S V L C D A E V> 43

160

GCT CTT GTT GTC TIC TCC CAT AAG GGA AAA CTC TTC GAA TAC TCC ACT
A L V V F S H K G K L F E Y S T> 59

220

GAT TCT TGT ATG GAG AAG ATA CTT GAA CGC TAT GAG AGG TAC TCT TAC
D S C M E K I L E R Y E R Y S Y> 75

GCC GAA AGA CAG CTT ATT GCA CCT GAG TCC GAC GTC AAT ACA AAC TGG
A E R Q L I A P E S D V N T N W> 91

280

TCG ATG GAG TAT AAC AGG CTT AAG GCT AAG ATT GAG CTT TTG GAG AGA
S M E Y N R L K A K I E L L E R> 107

340

AAC CAG AGG CAT TAT CTT GGG GAA GAC TTG CAA GCA ATG AGC CCT AAA
N Q R H Y L G E D L Q A M S P I> 123

400

GAG CTT CAG AAT CTG GAG CAG CAG CTT GAC ACT GCT CTT AAG CAC ATC
E L Q N L E Q Q L D T A L K H I> 139

460

GGC ACT AGA AAA AAC CAA CTT ATG TAC GAG TCC ATC AAT GAG CTC CAA
R T R K N Q L M Y E S I N E L Q> 155

AAA AAG GAG AAG GCC ATA CAG GAG CAA AAC AGC ATG CTT TCT AAA CAG
K K E K A I Q E Q N S M L S K Q> 171

FIG. IA

2 / 44

FIG 1B

3 / 44

TCTTAGAGGA AATAGTCCT TTAAAAGGGA TAAAA ATG GGA AGG GGT AGG GTT CAG	7
25	
TTG AAG AGG ATA GAA AAC AAG ATC AAT AGA CAA GTG ACA TTC TCG AAA	23
L K R I E N K I N R Q V T F S K	
85	
AGA AGA GCT GGT CTT ATG AAG AAA GCT CAT GAG ATC TCT GTT CTG TGT	39
R R A G L M K K A H E I S V L C	
145	
GAT GCT GAA GTT GCG CTT GTT GTC TTC TCC CAT AAG GGG AAA CTC TTT	55
D A E V A L V V F S H K G K L F	
205	
GAA TAC TCC ACT GAT TCT TGT ATG GAG AAG ATA CTT GAA CGC TAT GAG	71
E Y S T D S C M E K I L E R Y E	
AGA TAC TCT TAC GCC GAG AGA CAG CTT ATA GCA CCT GAG TCC GAC TCC	87
R Y S Y A E R Q L I A P E S D S	
265	
AAT ACG AAC TGG TCG ATG GAG TAT AAT AGG CTT AAG GCT AAG ATT GAG	103
N T N W S M E Y N R L K A K I E	
325	
CTT TTG GAG AGA AAC CAG AGG CAC TAT CTT GGG GAA GAC TTG CAA GCA	119
L L E R N Q R H Y L G E D L Q A	
385	
ATG AGC CCT AAG GAA CTC CAG AAT CTA GAG CAA CAG CTT GAT ACT GCT	135
M S P K E L Q N L E Q Q L D T A	
445	
CTT AAG CAC ATC CGC TCT AGA AAA AAC CAA CTT AGT TAC GAC TCC ATC	151
L K H I R S R K N Q L M Y D S I	
AAT GAG CTC CAA AGA AAG GAG AAA GCC ATA CAG GAA CAA AAC AGC ATG	167
N E L Q R K E K A I Q E Q N S M	
505	
CTT TCC AAG CAG ATT AAG GAG AGG GAA AAC GTT CTT AGG GCG CAA CAA	183
L S K Q I K E R E N V L R A Q Q	

FIG 2A

4 / 44

GAG CAA TGG GAC GAG CAG AAC CAT GGC CAT AAT ATG CCT CCG CCT CCA
 E Q W D E Q N H G H N M P P P P P 199
 CCC CCG CAG CAG CAT CAA ATC CAG CAT CCT TAC ATG CTC TCT CAT CAG
 P P Q Q H Q I Q H P Y M L S H Q 215
 CCA TCT CCT TTT CTC AAC ATG GGG GGG CTG TAT CAA GAA GAA GAT CAA
 P S P F L N M G G L Y Q E E D Q 231
 ATG GCA ATG AGG AGG AAC GAT CTC GAT CTG TCT CTT GAA CCC GGT TAT
 M A M R R N D L D L S L E P G Y 247
 AAC TGC AAT CTC GGC TGC
 N C N L G C 253

FIG. 2B

ATG GGA AGG GGT AGG GTT CAG TTG AAG AGG ATA GAA AAC AAG ATC AAT M G R G R V Q L K R I E N K I N	16
60	
AGA CAA GTG ACA TTC TCG AAA AGA AGA GCT GGT CTT ATG AAG AAA GCT R Q V T F S K R R A G L M K K A	32
120	
CAT GAG ATC TCT GTT CTG TGT GAT GCT GAA GTT GCG CTT GTT GTC TTC H E I S V L C D A E V A L V V F	48
180	
TCC CAT AAG GGG AAA CTC TTT GAA TAC CCC ACT GAT TCT TGT ATG GAG S H K G K L F E Y P T D S C M E	64
240	
GAG ATA CTT GAA CGC TAT GAG AGA TAC TCT TAC GCC GAG AGA CAG CTT E I L E R Y E R Y S Y A E R Q L	80
ATA GCA CCT GAG TCC GAC TCC AAT ACG AAC TGG TCG ATG GAG TAT AAT I A P E S D S N T N W S M E Y N	96
300	
AGG CTT AAG GCT AAG ATT GAG CTT TTG GAG AGA AAC CAG AGG CAC TAT R L K A K I E L L E R N Q R H Y	112
360	
CTT GGG GAA GAC TTG CAA GCA ATG AGC CCT AAG GAA CTC CAG AAT CTA L G E D L Q A M S P K E L Q N L	128
420	
GAG CAA CAG CTT GAT ACT GCT CTT AAG CAC ATC CGC TCT AGA AAA AAC E Q Q L D T A L K H I R S R K N	144
480	
CAA CTT ATG TAC GAC TCC ATC AAT GAG CTC CAA AGA AAG GAG AAA GCC Q L M Y D S I N E L Q R K E K A	160
ATA CAG GAA CAA AAC AGC ATG CTT TCC AAG CAG ATT AAG GAG AGG GAA I Q E Q N S M L S K Q I K E R E	176
540	
AAC GTT CTT AGG GCG CAA CAA GAG CAA TGG GAC GAG CAG AAC CAT GGC N V L R A Q Q E Q W D E Q N H G	192

FIG. 3A

6 / 44

600		
CAT AAT ATG CCT CCG CCT CCA CCC CCG CAG CAG CAT CAA ATC CAG CAT	H N M P P P P Q P Q Q H Q I Q H	208
660		
CCT TAC ATG CTC TCT CAT CAG CCA TCT CCT TTT CTC AAC ATG GGA GGG	P Y M L S H Q P S P F L N M G G	224
720		
CTG TAT CAA GAA GAA GAT CAA ATG GCA ATG AGG AGG AAC GAT CTC GAT	L Y Q E E D Q M A M R R N D L D	240
CTG TCT CTT GAA CCC GTT TAC AAC TGC AAC CTT GGC CGT CGC TGC TGA	L S L E P V Y N C N L G R R C	255

FIG. 3B

7 / 44

FIG. 4A

C	C	A	A	C	A	T	C	T	G	CCG	TTG	A	G	A	T	G	GGT	GAA	GAG	CTG	GCT	GGC	
P	H	N	I	C	F	P	P	L	P	M	D	R	G	E	E	L	A	L	A	R	C	A	240
GCG	GCG	GCG	GCG	CAG	CAG	CAG	CAG	Q	Q	CCA	CTG	CCG	GGG	CAG	GGG	CAA	CCG	CAA	CCG	CTC	CGC	ATC	780
A	A	A	A	Q	Q	Q	Q	P	P	P	L	P	G	Q	A	Q	P	Q	L	R	1	260	
GCA	GGT	CTG	CCA	CCA	TGG	ATG	CTG	AGC	CTC	AAT	GCA	TAA	GGAGAGGGTCGATGAACACATG	845									
A	G	L	P	P	W	M	M	S	H	N	A	*	273										
ACCTCCCTCTCTCTCGTCATGGATCATGACGTACGGTACCGTACCATATGGTTGCCTGCCCTCATCGATCG																							
CGAGCAATGGCACGCTCATGGATCATGGCTCCCTGGTTAACCCCTAGCCCTATGTTCATGGCGTCAGCAACT																							
AAGCTAACATTGTTATGCAAGAAAGGTAACCCGGCTAGGCTGTGATACTTGCCAGGATTCAGTATGCTTGT																							
TACTGCCAGTACCTTGAATCTAGGGCGTTACTTAAACATGGTGCAGTTACTTAAACATGGTGCATCT(A)n																							
TGTAATAGTAGTATTATCGATTGGGCATCT(A)n																							

FIG. 4B

9 / 44

T T A G G A A A T G G G A A G G G G T A G G G T T G A A T T G A G A G G A T A G A G A A C A A G
 M G R G R V E L K R I E N K > 14

51

A T C A A T A G A C A A G T G A C A T T C T C G A A A A G A A G A A C T G G T C T T G A G
 I N R Q V T F S K R R T G L L K > 30

111

A A A G C T C A G G A G A T C T C T G T G A T G C C G A G G T T T C C C T T A T T
 K A Q E I S V L C D A E V S L D > 46

171

G T C T T C T C A T A A G G C C A A A T T G T T C G A G T A C T C C T T C T G A A T C T T G C
 V F S H K G K L F E Y S S S E S C > 62

231

A T G G A G A A G G T A C T A G A C C G T A C G A G A G G T A T T C T T A C G C C G A G A G A
 M E K V L E R Y E R Y S Y A E R > 78

C A G C T G A T T G C A C C T G A C T C T C A C G T T A A T G C A C C A G A C C T C C T G A
 Q L I A P D S H V N A Q T N W S > 94

291

A T G G A G T A T A G C A G G C T T A A G G C C C A A G A T T G A G C T T T G G A G A G A A C
 M E Y S R L K A K I E L L E R N > 110

351

C A A A G G C A T T A T C T G G G A A G A G T T G G A A C C A A T G A G C C T C A A G G A T
 Q R H Y L G E E L E P M S L K D > 136

411

C T C C A A A T C T G G A C A G C A G C T T G A G A C T G C T C T T A A G C A C A T T C C C
 L Q N L E Q Q L E T A L K H I R > 152

471

T C C A G A A A A A T C A A C T C C A T G A A T G A G T C C C T C A A C C A C C T C C C A A A G A
 S R K N Q L M N E S L N H L Q R > 168

A A G G A G G G A G A T A C A G G A G G A A C A G C A T G C T T A C C A A A C A G A T A
 K E K E I Q E E N S M L T K Q D > 184

531

A A G G A G G G A A A C A T C C T A A A G A C A A A A C A A A C C C A A T G T G A G C A G
 K E R E N I L K T K Q T Q C E Q > 200

591

C T G A A C C C C A G G T C G A C G A T G T A C C A C A G C C A C A A C C A T T T C A A C A C

FIG 5A

10 / 44

L	N	R	S	V	D	D	V	P	Q	P	Q	P	F	Q	H>	216	
651																	
C	C	A	T	C	T	T	A	G	A	T	T	C	T	T	A	A	ATG
P	H	L	Y	M	I	A	H	Q	T	S	P	F	L	N	H>	232	
711																	
G	G	T	T	G	T	A	C	A	A	G	G	A	G	A	A	A	AAT
G	G	L	Y	Q	G	E	D	Q	T	A	M	R	R	N	H>	248	
*																	
C	T	G	G	A	T	C	T	G	A	T	T	G	T	T	A	C	GCC
L	D	L	T	L	E	P	I	Y	N	Y	L	G	C	Y	H>	262	
*																	
G	C	T	T	A	G	--										263	
A	*	X	H>														

FIG. 5B

ATG GGA AGG GGT AGG GTT GAA ATG AAG AGG ATA GAG AAC AAG ATC AAC M G R G R V E M K R I E N K I N	16
60	
CGA CAA GTG ACG TTT TCG AAA AGA AGA GCT GGT CTT TTG AAG AAA GCC R Q V T F S K R R A G L L K K K A	32
120	
CAT GAG ATC TCG ATC CTT TGT GAT GCT GAG GTT TCC CTT ATT GTC TTC H E I S I L C D A E V S L I V F	48
180	
TCC CAT AAG GGG AAA CTG TTC GAG TAC TCG TCT GAA TCT TGC ATG GAG S H K G K L F E Y S S E S C M E	64
240	
AAG GTA CTA GAA CAC TAC GAG AGG TAC TCT TAC GCC GAG AAA CAG CTA K V L E H Y E R Y S Y A E K Q L	80
300	
AAA GTT CCA GAC TCT CAC GTC AAT GCA CAA ACG AAC TGG TCA GTG GAA K V P D S H V N A Q T N W S V E	96
360	
TAT AGC AGG CTT AAG GCT AAG ATT GAG CTT TTG GAG AGA AAC CAA AGG Y S R L K A K I E L L E R N Q R	112
420	
CAT TAT CTG GGC GAA GAT TTA GAA TCA ATC AGC ATA AAG GAG CTA CAG H Y L G E D L E S I S I K E L Q	128
480	
AAT CTG GAG CAG CAG CTT GAC ACT TCT CTT AAA CAT ATT CGC TCG AGA N L E Q Q L D T S L K H I R S R	144
540	
AAA AAT CAA CTA ATG CAC GAG TCC CTC AAC CAC CTC CAA AGA AAG GAG K N Q L M H E S L N H L Q R K E	160
600	
AAA GAA ATA CTG GAG GAA AAC AGC ATG CTT GCC AAA CAG ATA AGG GAG K E I L E E N S M L A K Q I R E	176
660	
AGG GAG AGT ATC CTA AGG ACA CAT CAA AAC CAA TCA GAG CAG CAA AAC R E S I L R T H Q N Q S E Q Q N	192

FIG. 6A

12 / 44

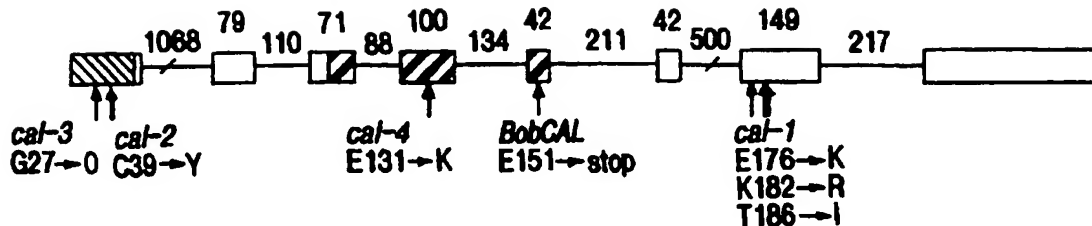
FIG. 6B

13/44

ATG GGA AGG GGT AGG GTT GAA ATG AAG AGG ATA GAG AAC AAG ATC AAC M G R G R V E M K R I E N K I N	16
60	
AGA CAA GTG ACG TTT TCG AAA AGA AGA GCT GGT CTT TTG AAG AAA GCC R Q V T F S K R R A G L L K K A	32
120	
CAT GAG ATC TCG ATT CTT TGT GAT GCT GAG GTT TCC CTT ATT GTC TTC H E I S I L C D A E V S L I V F	48
180	
TCC CAT AAG GGG AAA CTG TTC GAG TAC TCG TCT GAA TCT TGC ATG GAG S H K G K L F E Y S S E S C M E	64
240	
AAG GTA CTA GAA CGC TAC GAG AGG TAC TCT TAC GCC GAG AAA CAG CTA K V L E R Y E R Y S Y A E K Q L	80
300	
AAA GCT CCA GAC TCT CAC GTC AAT GCA CAA ACG AAC TGG TCA ATG GAA K A P D S H V N A Q T N W S M E	96
360	
TAT AGC AGG CTT AAG GCT AAG ATT GAG CTT TGG GAG AGG AAC CAA AGG Y S R L K A K I E L W E R N Q R	112
420	
CAT TAT CTG GGA GAA GAT TTA GAA TCA ATC AGC ATA AAG GAG CTA CAG H Y L G E D L E S I S I K E L Q	128
480	
AAT CTG GAG CAG CAG CTT GAC ACT TCT CTT AAA CAT H ATT CGC TCC AGA N L E Q Q L D T S L K H I R S R	144
540	
AAA AAT CAA CTA ATG CAC TAG T CCCTCA ACCACCTCCA AAGAAAGGAG K N Q L M H X	150
600	
CTAAGGACAC ATCAAAACCA ATCAGAGCAG CAAAACCGCA GCCACCATGT AGCTCCTCAG	
660	
CCGCAACCGC AGTTAAATCC TTACATGGCA TCATCTCCTT TCCTAAATAT GGGTGGCATG	
720	
TACCAAGGAG AATATCCAAC GGCGGTGAGG AGGAACCGTC TCGATCTGAC TCTTGAACCC	
ATTACAACT GCAACCTTGG TTACTTGCC GCATGA	

FIG. 7**SUBSTITUTE SHEET (RULE 26)**

14 / 44

**FIG. 8A**

CAL	MGRGRVLLKRIKNKINRQVTFSKRRTGLLKKAQKISVLCDAKVSLIVFSK	50
BoCAL	M A H I	
BobCAL	M A H I	
AP1	Q A H	A V
CAL	KGKLFEYSSESMEKVLERYERYSYAERQLIAPDSHVNAQTNW <u>SMEYSRL</u>	100
BoCAL	H K KV	
BobCAL	K K	
AP1	TD I E D -- N	
CAL	<u>KAKIELLERNORHYLGEELFPM</u> SLKD L ONLEQQLETALKHIRSRKNQLMY	150
BoCAL	D S I E D S	H
BobCAL	W D S I E D	T Y
AP1	D Q A P E D T	
CAL	<u>ESLNHLQRKEKEI</u> QEENSM L TKQIKERENILKT K QTQCEQLNRSVDDVPQ	200
BoCAL	V A R S R H N S Q HHVA	
BobCAL	*	
AP1	I E K A Q S K RAQ E WD Q QGHNMP -	
CAL	PQP <small>F</small> QHPHL---YMIAHQ <small>T</small> SPFLNMGGLYQGEDQTAMRRNNLDLTLEPIY	247
BoCAL	QLN YM -----AS M YP V R	
BobCAL		
AP1	L P QHQIQHP LS P ED PM D E V	
CAL	NY-LGCYAA*	255
BoCAL	CN YF	
BobCAL		
AP1	CN F	

FIG 8B

AAGCCATCTCTTAAAGGTAAACGAGGTTGGGATGAGGAGGAA	100
F R W N P T R A L V Q A P P P V P P L Q Q Q P V T P Q T A A F G	10
ATGCCACTTGCTGTTAGGGAACTTTCGGTCAACCCACCTCCGGT	200
N R L G G L E G L P G Y T A K I A S T L V	43
TGGGTATGAAGGACGGAGCTTGAAGAGATGTGATCTCATATCTT	300
G M K D E E L E S H M N S L S H I F R W E L L V G E R Y G I K A A	77
CCTAGAGCTGAACGGGACGGATTCAAGGAGGAACTTCTAGACGCCGT	400
V R A E R R L Q E E B S S R R H L L S A A G D S G T H	110
CACGGCTTGTATGCTCTCCAAAGAAATGATGGACAGGGTTATCTGAGGA	500
H A L D A L S Q E D D W T G L S E B P V Q Q Q D Q T D A A G N N G G	143
GAGGAGGAATGGTACTGGGACGGAGCTCAGGAGGAAAGGAGGAA	600
G G S G Y W D A G Q K M K Q Q Q R R R K P M L T S V E T D	177
CGAAGACGTAAACGAGGTGAGGATGACGAGGAAACGGGATTCGGTT	700
E D V N E G E D D D G H D N G N G G S G L G T E R Q R E H P P F I V	210
15/4	243

SUBSTITUTE SHEET (RULE 26)

ACGGAACCTGGGAAAGTGGCACGTGGCAACAGAACCCCTTAGATTATCTGTTCCACTTGACGAAACATGCCGTGAGTCCTTCTAGGTCAAGACAA
 T E P G E V A R G K N G L D Y L F H L Y E Q C R E F L L Q V O T I 277
 TTGCTAAAGACCGTGGGAAAAAATGCCCAACCAAGTGCAGTACGGTAAGGAAATCAGGGAGGTACATAAACAGCTTAAAT 1000
 A K D R G E K C P T K V T N Q V F R Y A K K S G A S Y I N K P K M 310
 GCGACACTACGTTCACTGTACGGTCTCCACTGCCTAGACGAAAGCTCAATTGCTCTAGAAGAGCGCTTAAGAGACCGGTGAGAACGTTGGCTCA
 R H Y V H C Y A L H C L D E E A S N A L R R A F K E R G E N V G S 343
 TGGCGTCAAGGCTTGTACAGGCCACTTGTAAACATGCCCTGCGTCATGGCATATAGACCCGCTTAAACGCTCATCTCTCTATTGGT
 W R Q A C Y K P L V N I A C R H G W D I D A V P N A H P R L S I W Y 377 / 44
 ATGTTCCAACAGCTACGTTGCTTGGCAATTGGGGAAACAATGGCGCTGGCTTACGTTGGCTTAACTTAGCTGTACCGATCGTC
 V P T K L R Q L C H L E R N N A V A A A L V G G I S C T G S S 410
 GACGTCTGGACCTGGTGGATCCGGGGACACTGGCTTCTAGTTGGCTTGGTGTATCGTTAGCTTAACATTAGTC
 T S G R G C G C D D L R F Stop 1400
 TTTAATTAGTCCTCTGGCTTAATTTCCTTAAACCTTAAATTGGTTATACAGCAGTTCTTAATGGCTTA 1500

FIG. 9B

17 / 44

GAATTCCCCG GATCTCCATA TACATATCAT ACATATATAT AGTATACTAT
 60
 CTTTAGACTG ATTTCCTCTAT ACACTATCTT TTAACTTATG TATCGTTCA
 120
 AAACTCAGGA CGTACATGGT TTAAATTGG TTATATAACC ACGACCATT
 180
 CAAGTATATA TGTCATACCA TACCGATTT AATATAACTT CTATGAAGAA
 240
 AATACATAAA GTGGGATTA AATGCAAGTG ACATCTTTT AGCATAGGT
 300
 CATTTGGCAT AGAAGAAATA TATAACTAAA AATGAACCTT AACTTAATA
 GATTTTACTA TATTACAATT TTTCCTTTTA CATGGCTAA TTATTTTC
 360
 TAAAATTAGT ATGATTGTTG TTTTGTGAA ACAATAATAC CGTAAGCAAT
 420
 AGTTGCTAA AGATGTCCAA ATATTTATAA ATTACAAAGT AAATCAATA
 480
 AGGAAGAAGA CACGTGGAAA ACACCAAATA AGAGAAGAAA TGGAAAAAAC
 540
 AGAAAGAAAT TTTTAACAA GAAAATCAA TTAGTCCTCA AACCTGAGAT
 600
 ATTTAAAGTA ATCAACTAAA ACAGGAAACAC TTGACTAACA AAGAAATTIG
 AAAATGTTGC CAACTTTCAC TTAAATTATAT TATTTCTCT AAGGCTTATG
 660
 CAAATATAGC CTTAAGCAAA TGGCGGAATCT GTTTTTTTT TTTGTTATG
 720
 GATATTGACT GAAATAAGG GGTTTTTC AACTTGAAGA TCTCAAAAGA
 780
 GAAAACATT ACAACGGAAA TTCATTGTA AAGAAGTGTAT TAAGCAAATT
 840

FIG 10A
 SUBSTITUTE SHEET (RULE 26)

18 / 44

GACCAAGGT TTTATGIGG TTATTCAT TATAATTG ACATCAAATT
 * * * * * 900
 GATAATATAT GGTGTTTA TTTAACATA TATAATTGATA TAACGTACAA
 * * * * *
 ACTAAATATG TTGATTGAC GAAAAAAAT ATATGTATGT TTGATTAACA
 * * * * * 960
 ACATAGCACA TATCAACTGA TTTTGACCT GATCATCTAC AACTAATAA
 * * * * * 1020
 GACACACAA CATTGAAAAA ATCTTGTACA AAATACTATT TTGGGTTTG
 * * * * * 1080
 AAAATTTGAA TACTTACAAT TATCTCTCG ATCTTCCCTCT CTTTCCCTAA
 * * * * * 1140
 ATCCGGCTA CAAATCGTC GACCCAAAC ATTACACAGT TGTCATTGG
 * * * * * 1200
 TTCTCAGCTC TACCAAAAC ATCTATTGCC AAAAGAAAAGG TCTATTGTA
 * * * * *
 CTTCACCTGTT ACACCTGAGA ACATTAATAA TAATAACCA AATTGATAAA
 * * * * * 1260
 ACAAAAGGGTT CTACACCTTAT TCCAAAAGAA TAGTGTAAA TAGGGTAAATA
 * * * * * 1320
 GAGAAATGTT AATAAAAGGA AATTAAAAAT AGATATTTTG GTGGGGTCA
 * * * * * 1380
 GATTTGTGTT CGTAGATCTA CAGGGAAATC TCCGGCGTC ATGCAAAGGC
 * * * * * 1440
 AAGGTGACAC TTGGGGAAAGG ACCAGTGGTC GTCACATGTT ACTTACCCAT
 * * * * * 1500
 TTCTCTTCAC GAGACGTCGA TAATCAAATT GTTATTTTC ATATTTTAA
 * * * * *
 GTCCCGAGTT TTATTAATAA ATCATGGACC CGACATTAGT ACCAGATATA
 * * * * * 1560
 CCAATGAGAA GTCGACACGG AAATCCTAAA GAAACCACTG TGGTTTTGC
 * * * * * 1620
 AACACAGAGA AACCAGCTTT AGCTTTCCC TAAAACCACT CTTACCCAAA

FIG. 10B

SUBSTITUTE SHEET (RULE 26)

19 / 44
1680

TCTCTCCATA AATAAAGATC CGGAGACTCA AACACAAAGTC TTTTTATAAA

1740

GGAAAGAAAG AAAAACTTTC CTAATTGGTT CATAACRAAG TCTGAGCTCT

1800

TCTTTATATC TCTCTTGAG TTTCTTATTG GGGGTCTTTC TTTTGTTCGG

TCTTTTACA GTAAGAAGTT TCTTAAAAAA GGATCAAAAA TGGGAAGGGG

1860

TAGGGTCAA TTGAAGAGGA TAGAGAACAA GATCAATAGA CAAGTGACAT

1920

TCTOGAAAAG AAGAGCTGGT CTTTGTAGA AAGCTCATGA GATCTCTGTT

1980

CCTCTGAGATG CTGAAGTTCGC CCTCTGAGAC TCTCTCCATA AGGGGAAACT

2040

CTTGAATAAC TCCACTGATT CTGGTAACT TCAACTAATT CTTACTTTT

2100

AAAAAAATCT TTAATCTGC TACTTATAT AGTTTTTTTC CCCC----GG

TCTATGATTG ATACTGTTT GTTATTATAA AGGTATCATA GAGATCGGT

2160

CTTGTGTTGT TTAGGAAAT CTGGTTAA TTGCATAAAA CCATCATTTAG

2220

ATTTATCTA AAATGTGATG ATATTTGGT CACATCTCCA TATTATTTAT

2280

ATATAAAAT GATAATTGGT TGATGATAAA GCTAACCCCA ATTCTGTGAA

2340

ATGATCAGTA TGGAGAAAGT ACTTGAACGC TATGAGAGGT ACTCTAACGC

2400

CGAAAGACAG CTTATGCCAC CTGAGTCGCA CGTCAATGTA TTTCAATAAA

TATTCTCCT TTTAATCCAC ATATATATTA TATCAATCTA TTTGTAGTAT

2460

FIG. 10C

SUBSTITUTE SHEET (RULE 26)

20/44

TGATGAATT TTTTGTATAA AAACCTTCGG TACACAGACA AACTGGTOGA
 2520
 TGGGTATTA CAGGCTTAAG GCTTAAGATTG AGCTTTGGA GAGAAACCA
 2580
 AGGTACACAT TTACACTCAT CACATTCTA TCTAGAAAAT CGATCGGTT
 2640
 CCATTTAAA GTAGTTAAA ATTCATGAT GCTATTGAA TTCAGGCATT
 2700
 ATCTTGGGGAA AGACTTGCAA GCAATGAGCC CTAAAGAGCT TCAGAACCTG
 * * * * *
 GAGCAGCAGC TTGACACTGC TCTTAAGCAC ATCCGACTA GAAAAGTATT
 2760
 * * * * *
 GCCTTCCT ATTTCGTTGA ACATATCTAT ATAATTTAA CCTTCACAAG
 2820
 * * * * *
 TGTTATTAAT ATGTGAACAT TGAAATACAT ATGTGTATGT ATCAATATAT
 2880
 * * * * *
 ATATCAGAA TCAATATCAA TTIGATATGT CTAAAGGTG GTTCGAATGT
 2940
 * * * * *
 ATGAGTTAAG TTGTTATTT TAAGACTCCA TATTACTAA AGTAATGGGT
 3000
 * * * * *
 TGTTATGT GATGIGIGTG TATCCAGAAC CAACTTATGT ACCAGTCAT
 * * * * *
 CAATGAGCTC CAAAAAAAGG TAATGAAAAC CCCTATCAA TGTTATGCTT
 3060
 * * * * *
 ATAGAGAAAC GTTCTGGAA GCTAATTAAC AATCGTGCGG TTTCGGAAAG
 3120
 * * * * *
 ACAGGAGAAAG CCCTTACAGG ACCAAAACAG CATGCCCTCT AACAGGAAC
 3180
 * * * * *
 ACATGTCATC ATTCCTCTT CATCAACATG TTGTCCTTG CATTACTGT
 3240
 * * * * *
 ACCTTCCACT GTTCTGCTCC ACACCTCCAG CCAGCTATA CCTACGGATAT
 3300
 * * * * *
 CTTCATATCT CCACCTTAAC TCGGCRCCAT TAAATAAAAA TAGAAAATCT

FIG. IOD
 SUBSTITUTE SHEET (RULE 26)

21 / 44

TTGCAAATTT GTTGAATA GCATAGATG TGTCATTG A TGTATAAT
 3360
 CACCA GCGTG TACCTAGATA TGGTTTGTCG GTTGTGTTT AAGGTGTCTC
 3420
 TGGATTGAA AATATTTTGA AATCTTTG AATGTTGTC CCATCATCT
 3480
 TACTTAGCTC ATATCTAATG ATATGAATAT AGACACTACT CCTATATATA
 3540
 AAATGTATA. ATAGTTCACTT GCATGAGTGC AACTGTGAAA ATAACCTATTT
 3600
 GTAAACCATG CATAATATATA GTTTCTCAC TTGAAATT GATGATGATA
 ATATGGTTG AAATAAATTT GCTGGCAGAT CAAGGAGAGG GAAAAAATTC
 3660
 TTAGGGCTCA ACAGGAGCAG TGGGATCAGC AGAACCAAGG CCACAATATG
 3720
 CCTCCCCCTC TGCCACCGCA GCAGCACCAA ATCCAGCAGC CTTACATGCT
 3780
 CTCCTCATCAG CCCTCTCTTT TTCTCAACAT GGGTAACAA AAAATCTCA
 3840
 ATCAGCTTA ATTAAAGCA CATACTTAT GCAAGCTAGT TACGTTAGGT
 3900
 GTTGTAAATT CATTGAAGTT ATAGCTGTTA GTGATGGTTA CATGATGCTA
 GATTTGAA CTAGAAAACT TTATTTAAA ACATTATTT ATTAAACGTAG
 3960
 GTTAAGCAA TGGTCCCAA ACGAACAAAC TTATCTAGT GTAAAAATGT
 4020
 ACATGGAAATG GTTGGAAAAA GCCTAAGTCG ACTTTTGTTG TTGTTGGCT
 4080
 ATGTGTAA GTACAATTT AGTTTGTTAG ATAAATGAA TTATATATC

4140

FIG. 10E

SUBSTITUTE SHEET (RULE 26)

22 / 44

TTTGACATTT CACAATGGAC TGATATTGGA TTTTCCCTTG TTGTACGGTG
4200
AAACATATGA TTACATATGC ACTTTCTAT ATATCCTATG TATGATTGTG
AATGCAGTGG TCGTGTATCAA GAAGATGATC CAATGGCAAT GAGGAGGAAT
4260
GATCTCGAAC TGACTCTTGA ACCCGTTTAC AACTGCAACC TTGGCCGTC
4320
GCCCCATGAA GCATTTCCAT ATATATATAT TTGTAAATCGT CACACATAAA
AACTAGTTTG CCATCATACA TATAAATAG

FIG. 10F

23 / 44

GCACCTGAGT CGGACTCCAA TGTAAACCAA TTTCCTCCAA TAACTTATA
 60
 TAAATTAAT ATTATTTCAAG TATTAGTGAT ATATACCTAT CTGTATTTAA
 120
 CTTGTGAGAT ATGGACCGAAC TGGTGGATGG AGTATAATAG GCTTAAGCT
 180
 AAGATTGAGC TTTTGGAGAG AAACCAAGGG TACATTTCA TCTCTCTTT
 240
 ATATTAATAG ATGAAATATC AAACAGGATT AATGTTAGTT AAAATGCCAT
 300
 GATTACTTAT AAGAAAATGA TGCATTTAA TAACTAAAAA ATGCATCGAT
 GCCTCATGAA AATTTAGGCA CTATCTGGG GAAGACTTGC AAGCAATGAG
 360
 CCCTTAAAGGA CTCCAGAAATC TAGAGCAACA GCTTGATCT GCTCTTAAGC
 420
 ACATCCGTC TAGAAAAGTA TGAATCTCC TATTCTTCA ATTAACATGT
 480
 ATACAACCTA AACACATATT ATTTTCTTAT TCAATACATA TATATGAATA
 540
 GTACATATGT GATTTCTTCG GTTGGTATAA AAGATCAAT CACGTCGATT
 600
 AGATGTATGA CTTTTTAAAG AATTAAGTATA TAGACTATGA TTAGTCATG
 TAATGGTACG TACGTTTATG CAGAACCAC TTAATGACGA CTCCATCAT
 660
 GAGCTCCAAA GAAAGGTATG TATAAACCT ATCAAATTGA CGTTTACATA
 720
 GATAACTGC GTGTAAAGAAT CCTATAGGGG AGCTAACAT CGTGGCGTTT
 780
 TGGAAATGAC AGGAGAAAGC CATAACGGAA CAAAACAGCA TGCTTTCCAA
 840

FIG. II A
SUBSTITUTE SHEET (RULE 26)

FIG. II B

SUBSTITUTE SHEET (RULE 26)

25 / 44

1680
ATAAAATAAAC TATCTTGTAT ATGGCCCTTTA CCAATTTCAC TACAAAACAT

1740
GTGATATTCTT CAGCACCTAT GTAGATAATT TGTAAGCTAT ATCATGTGCA

1800
TATGAATGTA AATGCAGGGG CCTTGATCAA GAAGAAGATC AAATGGCAAT

GAGGAGGAAC GATCTCGATC TGTCTCTTGA ACCCGGTTAC AACTGGCAACC

1860
TTGGCCCGTGG CGCGT

FIG. IIc

26 / 44

GAGCTCTTCT TTTATCTCT TCTTGTAGTT TCTTGTTCG TTGGTTTC
 60
 TTAGAGGAAA TAGTTCCTT AAAAGGGATA AAAATGGAA CGGGTAGGGT
 120
 TCAGTTGAAG AGCTTAGAAA ACAAGATCAA TAGACAACTG ACATTCCTCA
 180
 AAAGAAGAGC TGTCCTTAIG AAGAAAGCTC ATGAGATCTC TGTTCTGT
 240
 GATGCTGAG TTGGCTTGT TGTCTCTCC CATAAGGGGA AACTCTTGA
 300
 ATACCCACT GATTCTTGGT AACTTCTCA TTTAAGAAAC AAAA---TAC
 CCTAAGATTG TATTTACAT GATCATTTAC TTTTTTACA CAGTATATAC
 360
 TCTATGTATA TAATATGATC ATAATTGTT GATGATAAGA AGCTAGCCCT
 420
 AATTCTGTGA ATTGAACAGT ATGGAGGAGA TACTTGAACG CTATGAGAGA
 480
 TACTCTTACG CGGAGAGACA GCCTATAGCA CCTGAGTCGG ACTCCAATGT
 540
 AAACCAATTG CTCTCCATTA ACTTATATAA ATTAATATT ATTTCAGTAT
 600
 TAGTGATATA TACTTATCTG TATTAACCTT GTGAGATATA GACGACTGG
 TCGATGGAGT ATAATAGGCT TAAGGCTAAG ATTGAGCTT TGGAGAGAAA
 660
 CCAGAGGTAC ATTTCATTC ATCATTTATA TATATGATGA AATATCAAAC
 720
 AGGATTAATG TTAGTTAAAATGATGATT ACTTATAAAA AAATGATGCA
 780
 TTAAATAAC AAAAATGCC ATGGATGCTC TATTGAAATT TAGGCACTAT
 840

FIG. 12A
 SUBSTITUTE SHEET (RULE 26)

27/44

CTTGGGERAG ACTTGCAAGC AATGAGCCCT AAGGAACCTCC AGAAATCTAGA
900
GCAACAGCTT GATACTGCCTC TTAAGCACAT CGGCTCTAGA AAAGTATGAA
TCTCTCTATT TCTTTAATTAA ACATGTATAC AACTTAAACA CATATTATT
960
TATTATTCAA ATACATATAT ATAAATAGTA CATACTGATC TTATTTGGTT
1020
GGATTTGAAA AGATCAATCA CGTGGATTAG AATGTATGAC TTITTTAAAGA
1080
ATTAGTATAT AGAGTATGAT TAGTCATTGT AATGGATCGT TTATGCAGAA
1140
CCAACCTATG TACGACTCCA TCAATGAGCT CCAAAGAAAG GTATGTATAA
1200
ACCCATATCAA ATTGACGTTT ACATAGAATA ACTGGGTGTA AGAACCTTAT
AGGGGAGCTA AAAATCGTGC CGTTTGGAA ATGACAGGAG AAAGCCATAC
1260
AGGAACAAAA CAGCATGCTT TCCAAGCAGG TGCCATTGTG CATTATTTT
1320
ATPTCGTCAA AATGTTTCTT ATTGTAGATC TGTTAGCTTC CACTGTTCTC
1380
ACCACACTTC AAGCCAAGCT ATACCTACCT ACCGACTAC --- CCTACATTT
1440
GATGCTATT ATATGTATAT CTATTTAGAA GTCGTGGCTT TGAAATTGAA
1500
TGATGATATG GTATGGTATA AGTTGGTAAC AAACCTGGTGT GTGAAATTGA
AACTTGTCAG ATTAAGGAGA GGGAAAAGT TCTTACGGCG CAACAAAGAGC
1560
AATGGGACGA GCAGAACCTT GGCCATATAA TGCCCTCGCC TCCACCCCCCG
1620
CAGCAGGCATC AAAATCGAGCA TCTTACGTC CGTGGTAAAC TGGCTT

FIG 12B

28 / 44

1680

TTTCTCAAC ATGGGGTAGT TAAAAATTGG TTCCCTTCAC TTCAAGTAC

1740

ATATGTGTTA TATATACAAAG ATAGTTAGGT GTTATAAGTC CAGTGAGTTA

1800

AGTGTGTTA GTGATGGTTA GATGTCTAAA TTGTGAATA CAAGTACTAA

1860

GATTTTCAT GTATATATTT AAACGTATTA ATCATCAATC AAATGGTGT

1920

AAAAGAAACA GACTTATATT TTGGGGAAAA GTAGATGGAA TGGCTGCTAA

1980

AAGTCTAAGA AACCTTGGG ACCAGGTGTT TTTTATGTT GTCAAATTA

2040

AACTTGAGGT AGTTAGATAA ATAAACTATC TTGATATGG GCCTTTACCA

2100

ATTCACAC AAAACATGIG ATATTTCAAG CACCTATGTA GATTAATTG

2160

AGAGCTAT CATGTGCATA TGAATGTAAA TGTAAGGGCC TGTATCAAGA

AGAAGATCAA ATGGCAATGA GGAGGAACGA TCTCGATCTG TCTCTTGAAC

CGTTTACAA CTGCAACCTT GGCGCGTCGCT GCTGA

FIG. I2C

29 / 44

GGATCCCTCC GGAAGCCTTA GATCAATGGT AGTTGIGGT ATTTCAGAT
 60
 CAGATTCTTT TGAAATCCA GTAACATAGT CTGGAAATAT GATTIGCTTC
 120
 TTGGTCACCG TTACTGCTTC TCGGTTGTC ATTTGGATT TTACGGACTT
 180
 TTGATCACTA TGATAATTTC TCTTTCTTA CGTCGAGATG TGCTGCTTT
 240
 TTCTAGATTG AATTTCTCAA TGTTGCTTIG ATCAGTAAAC CATTGATT
 300
 CTTTCCCTCA TTGATGGATC CAATTTCTTC GGGAGATAAA TAAGGAAAAA
 ATGGACTATT ATTTTGAA AATACAGGAG AAAAATTC TTAGAATAA
 360
 AAAGATTTT ATAGTGACCA TGAATTTTGT TGTTTTTTA AAAAGAAAAA
 420
 AAAACTCGAT TGGATTGGAT GACACATGAA AATTAACATT CAAATGGAT
 480
 CTTAGTTAAC AGATATTGCA TCCACCATAT AATAAAATAT CATAATTATG
 540
 TGATGGCA GGTTTGTGTT GGTCAAAATG TTATTTAT CACAATTEAA
 600
 TAACAGATCA TTACCAATT TGTTTTTGA TAATTTATGC CAACTTAGTA
 AATTCATCCA AAAAGTTGAA AAATATAGAT GTGAAATATG TTGACGGATA
 660
 TACAACACTC AAAACANTAT ACTCAAAAAA AAAAAAATT GAAAGGGCA
 720
 ACGATTCAA CATATATGCT AAATTTAAT AATGGACAAA GGAGGAAGTA
 780
 CTGCATATGT ACAGAAAATG TTGATAATGG AGAGCAGOGG ATAGTGCGC
 840

FIG. 13A
 SUBSTITUTE SHEET (RULE 26)

30 / 44

CAAGGGCAGG AGCTTTAGAT TCTTTTAGTT TGCTCTAAAT GTCTCTCTT
900
GGTACTTTTA ATTGCCTTAG TTGCTTCCTT CTTATCTCCA CATAAATAAA
TGGGGTAACC ATTTTCCTTC GTATCTTATT CGGATCTTGT GATCTAIGTA
960
CGTACTACAT GAATAATCG TGTCAATAA GTTATATCA TTGGTCCTG
1020
TTAAAGTGAT CATGGTGAT TAATCTATAA TACGTAGTTC TCTTAATTG
1080
TTCCCTAGAA TTCCATCAA GACAAATTGT AGCAAAAAGA AAAGTTGAGT
1140
ATATAATTG CTAGTAGTA CAAAAAAA CTTATGGTA ATTGTATTT
1200
TGGATATTC CTNATTAAC CCAAACCTCA AAATTAATT TCTTCCTGGT
TACCTTATA TCCAACGTA AATCTATTGA CTCAACAAAA TACACAGTIG
1260
TCMATTGAAG TTCAACTCTA CCAAGAAACA TCTATATGTA CTTCACTGTT
1320
CTTACCCCCG AGCAATTAAA ACCTCTATAA CTACTGGTT ACATTATAC
1380
ATTTTATTT ACAAAAATA TATATCAACA ACCAATAATA TAGTTAGAAA
1440
ATGAAAGAAA ATTATTTAAG AAATATCGC CGTCAATGCA AATGAAATGC
1500
GACACTTGGG GAAGCTCTGA AGTCTGTGGT CTGTCATAT TTCACTTGT
TAGCTAACCC ATTTTCAGT CACTAGACGT CGATAATCAA TTATTGGTAT
1560
TTTTTTATC AATGTTCCAC TTATTGAAAA TTATATACGA GAAAACATAG
1620
ACTGACACATT AGGCAATGGA AGTCTAATCA GACCAATGAG AAGTCGACAA

FIG 13B

SUBSTITUTE SHEET (RULE 26)

31 / 44

1680

CACATCCTAG AAACCAACTC TGGTTTATTTC CCTTCCTCAA TACCAAGTTA

1740

TAGNNTTCTT TCAAACCGCT ATTTCCAAA TATCTCTCTT TTAAATTAAG

1800

AGTGAAAGAA GCACTCTTTC ACATTACCAT CATTAGAAAA CTTTCCTTAT

TAGATCAAGA TCGTGTAT CTCTCTGTT TTTCTCTCAT ATAATTTAGT

1860

TATTTTAA GAATGGGAA GGGGTAGGGT TGAATTGAAG AGGATAGAGA

1920

ACAAGATCAA TAGACAAGTG ACATTCTCGA AAAGAAGAAC TGGCTTTTG

1980

AAGAAAGCTC AGGAGATCTC TGTCTTTGT GATGCCGAGG TTTCCTTAT

2040

TGCTCTCTCC CATAAGGGCA AATTGTCGA GTACTCCCT GTAACTTGGT

2100

AATTGCTTAA TTCCCTCTTT TTTAATGTT ATTTTATGTC TGCCCTTCGGT

TGCCCTRACT AGTAGTCTTT GTTCTACTTA AGGCATATTT TCAGTGCTT

2160

CTATGCTTAA ATCTGTCTTT GCTGAAATTG TCCCCTGAT TTGGTATCTA

2220

TTTACTTGGG ATCTACGAAC TGATTGTTGGT GGTCATATCA TTAGTTTATT

2280

TTTATCAATA ATTATTTATA TATCAAAGAA AATGAAATTG TTTAGGACTT

2340

TTAGTGAACC CTACAAATCC ATCTACTTAA TTATAGTGGC ATGGATTTGT

2400

AAGAAATCTT CAGCATCTTC TTAAATCTGG AAATGTACAT TTTCCTCAA

GTCAAGTTA GTATATTAGG TACAGAAAGA ACCGATGTTT ATGGCTAGA

2460

FIG. 13C
SUBSTITUTE SHEET (RULE 26)

32 / 44

CTACGGTTTT TGCCTTCTAGG AAAGCTATAC TTTTCCTTAA ATATCTTCAA
 2520
 GTTGCATTTT ATGAACACAC ACACACATAT ATATATATAT ATATTAAGTAT
 2580
 ACCAATAATC TAAATTAAGT TTAGAAAGAA ACTCTTCATT TTTCCCATT
 2640
 TAATAATGGT TTATAGCTAG GTATAGAGAA ACTGGAATA AGTATGTGAC
 2700
 ATCTAAGTAT GGGGAGTCCT TGACCTCTGG GGATTAATGT AAAACAGATC
 * * * * *
 GTTCTTTTTT TTCTAAACAG TTCCCTCGTA CTGATGGTCA AACTTAACCT
 2760
 CAAACAGTTCC TTTTAAACCTT TTATAGGGTG CTGAAATACG TCTTGGGGTG
 2820
 TGGGGTTAGT GGCTCTACTG GTTTATTAT TTTTAAAAT GGTAGAAATC
 2880
 AGTACTGTTT CTAGCTAGGG TTTAGGCACA AAACTAGAGA TCATCTTAT
 2940
 TCCATTAATAG AAAGGAAAGAA ACTAATGTTT AATGACATAG ATTAATTAGA
 3000
 TAACCCTACA TAATCAGATG CTATATGTTA TCACATATTG TGGGTGAATC
 * * * * *
 GTTAATTACG TTGAAACAA GTGGCCTCTT GTGCTAGCTG ATAAGATAGT
 3060
 TGNGTATGCA ATTATATTGG TGGTGAATC CAAACTAATT CTAACTCGTA
 3120
 AGCTTAATAT TTGTAGCATG GAGAAGGTAC TAGAACGCTA CGAGAGGTAT
 3180
 TCTTACGCCG AGAGACAGCT GATTGCACT GACTCTCAOG TERAATGTATG
 3240
 TTTAATGGTC TCCATCATAT ATTTGCTAT ATTTGAAATC TTGCATGTGT
 3300
 TTTAACATAG CATATAACTG ATTATGGCT TTCAATGTTGG AAATTAAATG

FIG. 13D
 SUBSTITUTE SHEET (RULE 26)

33 / 44

TGAAGGCACA GACGAACTGG TCAATGGAGT ATAGCAGGCT TAAGGCCAAG
 3360
 ATTGAGCTTT TGGAGAGAAA CCRAAGGTAC ATAGTACATT TAAATTTATT
 3420
 GTAGTAGTTA AATATTGAGG AATAACAGAA GAGAGAAATGT TCTTAATTAA
 3480
 CTAATCTTC ATAGGCATTA TCTGGGAGAA GAGTTGGAAC CAATGAGCCT
 3540
 CAGGAACTTC CAAATCTGG AGCAGCAGCT TGAGACTGCT CTAAAGCACA
 3600
 TTGCTCCAG AAAAGTGCTGT AAATATATCC CACACCTAT CTCATGCTAT
 AACTAACTTT GACTTGTGT GGATGTATTA CATATAGTC AATATTGTAT
 3660
 AGAGATTGTC TCATATAAAT AAATAATTCTT TGGCCTTTT GTATGCAGAA
 3720
 TCAACTCATG AATGAGTOCC TCAACCACCT CCAAAGAAAG GTAGCTAAGT
 3780
 TAAACCATT TTATCTCTCA AGTCCCTGTGT GTATAGAGTC ATGACTTATA
 3840
 TGTTAGAGAT ATAATCTTT TAATAAATAA ATAACATATA GGTTATATAT
 3900
 AATTAGGTT AATATATTAT TAATTACTAG ATGTATATAT ACTTATATAG
 ATCTATATAA AAGAGAAATT GACAATGGTG TCATTTTGT GGAATGACA
 3960
 GGAGAACGAG ATACAGGAGG AAAACAGCAT GCTTACCAAACAGGTGATCA
 4020
 TGTTCCTTGTG CATTCTAAC TGTTCACTA TTTACANTTC CACTGTTGAA
 4080
 CTCCACTTCA ATCTCTACCT TAACGTACCA TCTCTCCACT TTGGCCCCA

FIG. 13E

4140

SUBSTITUTE SHEET (RULE 26)

34 / 44

ACTCTTTGCA GAAAAAGAA TIGATATGTA GTTTCCTTG ATGGTATAA
 * * * * * 4200
 TCATGAGCCT AGCTCCACGT ATAGGTAAAC C TTGTCGCCCT TAGTATTAAG
 * * * * *
 GTTGTCTCCC AGATTGAAC TTGAACCTGA ACTGTCTCT CATAATCATA
 * * * * * 4260
 GTCATGCT AAATTACACA TACATAGCT AGATAGCTAG GAGCTATATT
 * * * * * 4320
 TTAAGTTTA TTGAGAAGTA AGAAAACGTA CGATGAAACT ACTTGATTA
 * * * * * 4380
 GACATATAT TAAATGAAAA AATATCACAA TAGTAAGACC TTGACGACCC
 * * * * * 4440
 TMAATTCCG TTAACATTT GCAGATTAA TTATCTTTT GCATTTGTT
 * * * * * 4500
 TGAAAATATC ATATTACAAA AAAAAGTATA AGAATAAAA ATTGAAGTTC
 * * * * *
 CTGAAATAAA TGCAAATAGC TGATTTAGTTC CAAATGGAA TCTATATAAC
 * * * * * 4560
 GATGATGCTT ATATCATTTT CTGGCGTGT GAAATCGGTAA TAGATAAAGC
 * * * * * 4620
 AGAGGGAAAAA CATCCCTAACG ACAAAACAAA CCCAATGTGA GCAGCTGAAC
 * * * * * 4680
 OGCAGCGTCC ACCATGTACC ACAGCCACAA CCTTTCAAC ACCCCCCATCT
 * * * * * 4740
 TTACATGATC GCTCATCAGA CTCTCCCTT CCTAAATATG CGGTAAACGGC
 * * * * * 4800
 AGTATTTCTT ATTTTTTAA GTTCTTTTTT CTAAACATAA TGTCAAATTTC
 * * * * *
 TCATATATAG TGAAGTGTTC TCAGTCAGTC ATATAGGCAA TGATAGTGAA
 * * * * * 4860
 TGCACCTCAT ATATAGGGTT TGTGTAGGT ATGGCGTAG AGGTGATGG
 * * * * * 4920
 TATGCATGCA TATTATGTA TTATGATTTT TAATTTGCTA TATATGATTG

FIG. 13F

SUBSTITUTE SHEET (RULE 26)

35 / 44

4980

TAATTTCACT GGTGTTGACCC AAGGAGAAGA CCAAACGGCG ATGAGGAGGA

5040

ACAATCTGGA TCTGACTCTT GAACCCATT ACAATTACCT TGGCTGTTAC

5100

GCCGCTTGAA TAGACTACAT CGATCTATAT CAATCTCTTT AAAATAATAT

AAGATCGATC CTCTATTTCAT GATCTATATT AAACACCGGT TAATTAATAT

5160

ATTTTGGTA TGCTCTTAA TCATATCAC ACATCAAGC CTTTTCCAA

5220

TICAATATAT CTCTGTTTTTC GGGGACCAT GAACTAAATGT AAATTTTGTC

5280

GACTGAGAGA CCTAGAAAGA ATTGGTGTTC AAACCTTTTC TATATTGATC

5340

TCATCGTAC ATTGAAATTG GATTTCTTTC ACACCCAAA ATATTTGEAA

5400

TACGAATTAA GTCTTGTATG ATTGAACTT TACTGGTCA AAGTAAATCA

CAGCCTTAA AGGAAATTG TGAATTGAAA ATAGAAATAA AAAATGTGGG

5460

AACGTGACAT TCGGTTCTT CTCCTTTC TTCTATGTAGG TCGGTGATAC

5520

GATCGGAAT GAGAATTATT GGGCCCTTGT GGCCTTCATA ATTATTAGTT

5580

CTTGTGAAAC GCGATAATA CTTGGCATTT TTGCAAAGA AGAAACTGTA

5640

TAAAGAAAT CGGAGAAGAA AAGAAAATA CTAGTCGGGG CAATGGAGGA

5700

TCTATGGAAG AGGGCAAAAT CGTTGGCAGA AGAAGGGGT AAGAAGTCIC

AGACGATAAC ACAATCAATCC TCGGCGACCT TCGTCAAATCT CGTCACCGAG

5760

FIG 13G

36 / 44

TCTAGATAAT CTTCTCAAGA AGGATTITAGA ATGGCATAAT CCRAAGGCCTC
 60
 AAACTCTGGC ATCTGAAACC ATATTAATCAA TTATATCATG ATTTCAGGATC
 120
 CAACCAATTAA AAAATRATCA GTGCATATGA TTTCATTAAGT CTCTCGACCA
 180
 AAACACTTAA CTACTCGATC ATGGTGCGAA ACAAGTCGAG AATGCTAGGT
 240
 CTATAATGAGA TGCTTAGGCC ACACGGCAGC TAATGTGATA CAAOGATCT
 300
 AGAGATCGGT TCTGAGATAT GCAAGCAGG TCACACGACC ATTCAATATAT
 GGIGTCTCTC TAGGCCACAC GCACAGCTAT GATGCATTAA CCCACACGGC
 360
 TTCAATCAC ATGATGCAAC AATGTGATCT ATCAAGGG ---CTCGAGC
 420
 TGCACACAGA CGGACCGCGAG CTGGCTGTGG TCGGATGCGA GCTGAAACGGG
 480
 ACGGGACTCG TCTGCTTCCT ATCGGGTTGG CGAGCTGCTT CCTATCGGGT
 540
 TTCAAGCGG CTGATCGGGA TTACAAGCTG GTTGTGAGG AACACGGAGCT
 600
 GGCTGTGATG CGAACCGGAAG CTGAGGTGT CTAGGATCAG GAACACCTTA
 GGGATGGAGC TGATCGGTG CTGACGGAGCT GGAACCGCGAG CTAGGACGAA
 660
 TEAGGGTTGG TCGGGATTAG GTTAAAGTCG CGCGCTAGGT TAGGTTAAAG
 720
 GGATGGCGA TTTAGCTTA GATTCAGAG AACAAATCGTG CTGATAACAT
 780
 GTTGTAAATTAA GAAGATTGAA GATTGAATAG TTCTGTGTTT TATTAACATA
 840

FIG. 14A
 SUBSTITUTE SHEET (RULE 26)

ACATGAAATT- ——AAAGAT TCCACGGAGTT TCGTACATGT TCTATTGCTA
900
CTTAGGTTAA GGGAGTTAAG CAAAGTAGAG TGATTGGCAT TAATCTTCA
GTAATGCCCA CGAAGACTCT AGTTAGAAGT CAGTTCAANTC TGACAAGCTG
960
TTAGAGGTTT ACTAACACTT GAGTTGGAT CTTGAAGGC CTTATATAG
1020
TTAAACGTAG ACCCAATATA ATACAAACT ATAGTATTTA CTTAAATT
1080
GAGTGCTCAC ACCAACCTGT TTAAAGCAAGA CAGGTCCCGA GACGGAGTC
GTTACGTTGT TGAGGCTC---

FIG. 14B

38 / 44

AAGCTTTCAGG GTTTTAGGGT TTTTGATTCG AAGCTTTCAGG GTTTTCATAA
 60
 TTCAAGATCAG AACAAATCAAT CAAACATGTC TAATGGAATC GATTTCATC
 120
 TAGTGTTAT AAGATGATCA GTTTTAGGTT ATACCAAATT TTAGGATTCA
 180
 TCAAGATCAG TGGATTCCG TAATAATGGA TTGGGGTTT AGGGTTTGT
 240
 CATAATGTTT TTAGATTAAT CGGTATACTT TIGTTTGAG GGTGAAACC
 300
 GGACCACCAA AGAGAACCGG TGAAACCTCGA CCTGCCACACC GACAGATGCC
 * * * * *
 AGCTGGCTGT CGTGGATGCC GAGCTGMAAG GGACGGGAGG CGTCTGCTTC
 360
 CTATGGGTTT CGCGAGCTGC TTCCCATCGG GTTTCGAAGC GGCTGATCGG
 420
 * * * * *
 GATTGGGAGC TGGTTGATCC GGAACACGGAG CTGGCTGAGA TCGGAACCGG
 480
 * * * * *
 AGCTGAGGTC GTCAGGATC AGGAACACCT TACGGATGGA CCTKGATCGGT
 540
 * * * * *
 TGCTGACGGG CTGGAAOGCG AGCTAGGGACA AATTTAGGGTT CGTCCGGATT
 600
 * * * * *
 AGCTTAAAGT CGCCGGCTAG GTTACGGTTA AGGGATGCG GATTTTACGT
 * * * * *
 TAGATTCGAG AGAACAAATCG TCCCTGATAAC GTGTTGTAAA ACAAACGGTT
 660
 * * * * *
 TTAGAAACTG AATGTTTATG TGTATTAATTA ATCATAATAT GGGTTTTT-
 720
 -----T ACAGTGCGAG AATGATAGAC TCGCTATGCC AATGAGTCC
 780
 * * * * *
 AGTCAGACCA ATGAGAAAGTC GACAGCAAAA CCTAGTAAAC TACTCTGTT
 840
 * * * * *

FIG 15A

SUBSTITUTE SHEET (RULE 26)

39 / 44

TTATCCTTGT CCAAAACCGG CTTCAGGGTTT CCCCTGAAACC GCTTAATTCCA
900
AAACATCTTC TCCTTAAATA AAGAAAGACT CTTTACACATT GTTATATCTCA
TCAGAAGGGG AAGAAGAAAA ACTTTCCCAA TTAGATCGAG CTTGCGGTTA
960
TCCTCTCTATT ATAGTTTATA TTCTCTACTG GGGCTTGTGTT GGTTGCTTCT
1020
CTTTTGGAC TTCTTTATA TAATTTATAT ATTCTACGGAG AAATGGGAAG
1080
GGGTAGGGTT GAAATGAGA GGATAGAGAA CAGATCAAC AGACAAGTGA
1140
CGTTTTCGAA AAGAAGAGCT GGTCCTTTGA AGAAAGGCCA TGAGATCTCG
1200
ATTCCTTGTG ATGGTGGAGT TTOCCTTATT GTCTCTCCC ATAAGGGAA
ACTGTTGGAG TACTGGCTG AATCTGGTA ACTGCATAAT TCCCTTTTA
1260
ATGTTTCTAG TGTCCTTTG TTGCCCCCAA TAAATAGTTT TGTGCTCCT
1320
TTAGGCCATT TCTGGTATC TTCTTATGTT TTTATGAAAAA TTCTCACAAA
1380
TTTGTGAGTT AATTACTTGG ATCTACGAAT TGATTCACC AAAGTGAAAT
1440
TAAACCATT TAGCATATTT CCTTATATCA GAAGAAATAA AAAAATAG
1500
GGCATAATAA GGTGTTATGT GAAGTGAAAG TTCTCTCAG GTCACACGTT
ATTAAGATAT CCTTAACCT AGATCAAGAT CTACTTCTAC TGGTGGGAC
1560
ATGGATTAC AAGAAATCGT CACTGTATAT GAACTTTAAT TAAACATGT
1620
ATAGACCTTT TTGTTCAA TAGAGAGTTA AGTAAATTAAC TCAAGAAAG

FIG. 15B

40 / 44 1680
AACCAACGTT ATGTCATCT AGGCTAGAGT GATTTTGCC TAAACAATTT
1740
GAAAAGCTGT CCTTATGCTT AAATATCTTT CAGCAGCATA GTAGTATGAA
1800
AGAAAATATT TCAATATCGT TGTATAAAGG TTCTTATAATT TTGGTTTTT
1860
TTTTTTCCG AAAAGCTTAA TATAGAGAAA CTAGAAGCTAG GGATGTGACA
1920
TCTAGGTATA GGGGTCTTTC ACCTCTGGGA TCAATGTAAA AGAGACCATT
1980
CTATTTCTA TCAACTCTC AGTTTCCGAT GGTCAAACCT TAACCTCAC
2040
AACTGTCCCC CTTTCAGAA GAGGACAAAC TATTATATGT ATATTAAGTT
2100
ATGTCGTTTC ATACATAAAAT ATCTAATAAC AAATTTATTT TTAAAAACAT
2160
ATAACAAAC TTATTTGAAG AATTGGAAAC TCAAAACGGG GACATATAGG
ACGGCTGCACG TCTAGAGGTG TGGGGTTAGT GATTCAACGG GTTTTTAATG
2220
TAGAGAAACT GTAGATGTAA GATTTGTTCT AGGGTTAAGG CACTAAACCA
2280
GGGATTATCT CTTTCCATC ATAAAAGTTA ATGTCCTAAA TGCATCGCTA
2340
ATTAATTAGG CAAACTAGAT GATAGTACGT AGTGTGTG TGIGIGIGTA
2400
TGGGATTTT TGGGTTAATA GTTACATCTT AGACAAATGT GTGGTCTCT
2460
GATAAGCTGA GAAAATATTG GGGTGCAGAC TCTTAGTGGT AATTAATTAT
ATCTAGAAAN NCCCANATAC NAATTTAATA CGGCTACTTT TTGGGTGAAT

FIG 15C
SUBSTITUTE SHEET (RULE 26)

41 / 44
GAATCTACAC TAAACCTTAAG CCTTAATGATA GCATGGAGAA GGTACTAGAA

2520

CGCTAOGAGA GGTACTCTTA CGCGGAGAAA CAGCTAAAGC CTCCGAGACT

2580

TCAAGTCAT GCAAGTTAA TGATCTCCAA GACTCTGTCA AACATATATG

2640

TACTATATCT TGAATGTTT TTCTTAATTA ACATAATTGA TGCACGTGTT

2700

ACATATGAA AATTAATTGT GTAGGCACAA AGGAACTGGT CAATGGAATA

TAGCAGGCTT AAGGCTAAGA TTGAGCTTTC GGAGAGGAAC CAAGGTACT

2760

TATAGCATTT AGGAACTTGC ATGTTAAAT AATAGTTTAT TGATTTAGTT

2820

TTTTTGTAA AAATTATTGT ATTAGTTAA CACTGGGAAT TAACTAAAAA

2880

GATGGTGGA TGGATTAATC ATAGGCATTA TCTGGGAGAA GATTTAGAT

2940

CAATCAGCAT AAAGGAGCTA CAGAAATCTGG ACCACCACT TGACACTCT

3000

CTTAACATA TTCGCTCCAG AAAAGTGTGT AAATAAGCAC ATACAAACGC

AAACATCTCT ATCTTATCTT TGAGTTTGTG AAGATAATAA TGCCTAATTT

3060

TATATAGAGT TTGATCTATA TGAATGAATA CAATTGAAC TCAATTGTAT

3120

GCAGAACTAA CTAAATGCCT AGTCCCCTAA CCACCTCCAA AGAAAGGTAC

3180

GTAAAAACCA TTTCATCTCT CAAGTCGTAC GIGGTTAGT GAGACTTTC

3240

TTACGGTTA AATCTTTCTAG TAAATACAA AACATATGGT TTTACACATG

3300

TTAGACTATT TGGTGAAGG AAACATTTGA AATGTAACA AAGGGGTTT

FIG 15D

SUBSTITUTE SHEET (RULE 26)

42 / 44

TGGGATTGAA TAAAATTTAA CATTCACTCA AAAAAAACAT ATGGTTTCATA
 3360
 TATATATTCG GTTATATATGA TTATATATAT ATATTTATAT AGGTAAATAT
 3420
 ATTAGTGTTT AATTATATAGT GTCACATAT AGATGTAGAA AGAACCTCIA
 3480
 GAGCGATCCC TGAGAATTTGT TICATTTGT AAAATTGACA GGAGAAAGAA
 3540
 ATACIGGAGG AAAACAGCT GCTTGCCAA CAGGTAACTCA TGTGTTGTC
 3600
 CATTTTTTAC TGTTTACAA CTGTTTACT ATTAAACTC CACTGTTCTA
 CTCCACTTCA ACCTTAACCT ACCATTGCTC AACCTTGGC ACCAACTCTT
 3660
 TTTTAAAAAG GAAGAATTAG TGTTTTCATG TGATTGGTAT AATCATGAGC
 3720
 ATATGTGCAC ACATGTAGGT GGGCTTTGTC CGTTTAGTAT TAAGGGTGT
 3780
 TCCTAGAATT GAACTGAAAC TGTCCTCTCG TAATCATAGT CTATATATAA
 3840
 CACCGTGCAC ATACAGTAGC CAGTAGGTTT ATTGAGCAA GATAC-----
 3900
 ---TGCTCTT ACTGTAATAC CGTGCCAAACA TTGATTTGAA TTGGATAACAT
 * * * * *
 AAATTEAGTT GATCATAACG TTTATGGTA TTTGAAATTG GTAGATAAAG
 3960
 GAGACGGAGA GTATCCTAAG GACACATCAA AACCAATCAG ACCAGCAAAA
 4020
 CGCGAGGCCAC CTGCTAGCTC CTCAGCGCA ACCGGAGTAA AATCTTACA
 4080
 TGGCATCTC TCCCTTCCTA AATATGGGGT AACGGTAGTG TTTCACTTTT

4140

FIG 15E

43 / 44

ATCTTGGTACACATAATAC ATATAGATCC GACACTCTG GIGTTAGTAA
4200
TTCAGTGTAT CGGATGAJGT TGTATGTATG TATGTTCAATA TTGGGGTT
GTGTTAAGTG TGGCGTTAGA GGTGATGGC TTTGTAACTA CAGTCTAGA
4260
ACTATACAAAT ATTAATAAG ATGGAATGAT ATATATATAT ACATATATTT
4320
TATTTCGCCA TATGATTCGIG ATTTCAGTGG CATGTACCAA GGAGAATATC
4380
CAACGGGGT GAGGAGGAAC CGTCTCGATC TGACTCTTGA ACCCATTCAC
4440
AACTGCAACC TTGGTTACTT TGCGGCATGA ATGGACTOGC CATATATOGA
4500
CTTAAATAAA TTATATAAG ATCGATTTT ACCTATAATA ATAGGCAGCA
ATGGTCTGCC ACCATATCTA TATACACTGG AAATTCTATT TATC----TT
4560
ACATTGATTT ATACTACATA AACCCCTCCAG ACCAAACTCG TCTCCATGCC
4620
AACTGATAGA TTCTCTAGAC ATGCTACACA CTCCATGACT CGGACTTATT
4680
TTGGTTTGG CGTTTCTCT GTTTTCTTA ATTGTTTGA ATTCTACTCT
4740
TTCACGATAT TTAAATTTT TCAACTTAT TTMGTMGCT CACAGTGAA
4800
AAATCTTCG TGAAGAAGTC GTATATATTC TGIGGGACCA CTCCCCCAAT
GTCCTCTGGT GGATCC

FIG. 15F

T K K K I K G I Q Q A T A G V S C W T S E N P W X T I V P
 ACA AAC AAA ATC AAA GGG ATT CGG CAA GCC ACT GCA CGA GTC TCA CAA GAC ACT TCG GAA ATG CCT AAC AAA ACA ATA GTT CCT 540

A A L P Q L T P T L V S L E V I E P E V L Y A Q Y D S S V
 GCA GCA TTA CCA CGG CTC ACC CCT ACC TGG TCA CTC CTC CGG ATT GAA CCC GAG ATT GAA GCA GGA TAT GAT AGC TCT GTT 570

P D S A W R I M T I L N H L C G R U V I A A V X W A X A I L
 CCA GAT TCA GCA TGG AGA ATT ATG ACC ACA CTC AAC ATG TTA GGT CGG CGT CGA GTC ATT GCA GCA TGG AAA TGG GCA AGG GCG ATA CTA 600

G L R N L H L D D Q M T L L Q Y S W M F L H A F A L G W R S
 GGC TTG AGA AAC TTA CAC CTC GAT GAC CAA ATG ACC CTC ATA CGG TCA TGG ATG TTT CTC ATA GCA TTT GCC TTD GGT TGA AGA TCA 630

Y R Q S S G N L L C P A P D L I I N E Q R H S L P C M Y D Q
 TAC AGA CGA TCA AGC GGA AAC CTG CTC TCC TTT GCT CCT GAT CTC ATT ATT ATG GAG CAG AGA ATG TCT CTA CCC TGC ATG TAT GAC CAA 660 / 44

C K B M L L V S S I L Q R L Q V S X I E Y L C H Y T L L L
 TGT AAA CAC ATG CTG TTT GTC TCC TCC TCA AGA TTA CGG GTC TCC TAT GAA GAG TAT CTC TGT ATG AAA ACC TTA CGT CTC CTC 690 / 44

S S V P Y E Q L K S Q E L P D E I R H T Y I K E L G K A I V
 TCC TCA GTT GCT AGG GAA GGT CTC AAC AGG CCA GAG TTA TTT GAT GAG ATT CGA ATG ACT TAT ATC AAA GAG CTC GGA AAA GCC ATC GTC 720

Y R E G H S S Q N W Q R F Y Q L T K L L D S M H E V V E H L
 AAA AGG GAA GGG AAC TCC ATG CAG AAC TGG CAA CGG TTT TAC CAA CTC ACA AAC CCTT CTC AAC TCC ATG CAT GAG GTC ATG ATC CTC 750

L T Y C F Q T F L D Y T M S I E F P E M L A E I I T H Q I P
 CTT ACC TAC TGC TTC CAG ACA TTT TTT GAT AGG ACC ATG ACT ATT GAA TTC CCA CGG ATG TTA CCT GAA ATC ATC ACT AAT CGG ATA CCA 780

K Y S N G N I Y Y L L P H Q Y stop
 AAA TAT TCA ATT GGA ATT ATC AAA AGG CGG ATT GCA TTC CCA CGG ATG TTA CCT GAA ATC ATC ACT AAT CGG ATA CCA 780

FIG. 16

SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US96/01041

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) :C12N 5/04, 15/10, 15/29, 15/82; C12P 21/02, 21/08
US CL :435/6, 172.3, 240.4, 320.1; 530/300, 350; 536/23.6, 24.1

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 172.3, 240.4, 320.1; 530/300, 350; 536/23.6, 24.1

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	ANTHONY et al. Cloning and sequence analysis of a <u>flo/fly</u> homologue isolated from cauliflower (<i>Brassica oleracea</i> L. var. <u>botrytis</u>). Plant Molecular Biology. 1993, Vol. 22, No. 6, pages 1163-1166, especially page 1164.	1-19
Y	ANTHONY et al. The cDNA Sequence of a Cauliflower <u>apetala-1/squamosa</u> Homolog. Plant Physiology. 1995, Vol. 108, No. 1, pages 441-442, especially page 441.	1-19
Y	CHUNG et al. Early flowering and reduced apical dominance result from ectopic expression of a rice MADS box gene. Plant Molecular Biology. October 1994, Vol. 26, No. 2, pages 657-665, especially page 657.	1-19

Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents:	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
*'A'		document defining the general state of the art which is not considered to be of particular relevance
*'E'		earlier document published on or after the international filing date
*'L'		document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
*'O'		document referring to an oral disclosure, use, exhibition or other means
*'P'		document published prior to the international filing date but later than the priority date claimed
"X"		document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"Y"		document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"&"		document member of the same patent family

Date of the actual completion of the international search

13 MAY 1996

Date of mailing of the international search report

31 MAY 1996

Name and mailing address of the ISA/US
Commissioner of Patents and Trademarks
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/01041

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	SOMMER et al. <u>Deficiens</u> , a homeotic gene involved in the control of flower morphogenesis in <u>Antirrhinum majus</u> : the protein shows homology to transcription factors. The EMBO Journal. 1990, Vol. 9, No. 3, pages 605-613, especially pages 609-610.	20-22
Y	SCOTT et al. Molecular and cellular aspects of plant reproduction. Cambridge, Great Britain: Cambridge University Press. 1994, pages 18-29, especially pages 21-22.	25
Y	KEMPIN et al. Molecular Basis of the <u>cauliflower</u> Phenotype in <u>Arabidopsis</u> . Science. 27 January 1995, Vol. 267, pages 522-525, especially pages 522 and 524.	27-31
Y	HULBERT et al. Recombination at the <u>Rp1</u> locus of maize. Molecular and Cellular Genetics. 1991, Vol. 226, pages 377-382, especially page 377.	27-31

INTERNATIONAL SEARCH REPORTInternational application No.
PCT/US96/01041**Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)**

This international report has not been established in respect of certain claims under Article 17(3)(a) for the following reasons:

1. Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:

2. Claims Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

Please See Extra Sheet.

1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
1-22, 25 and 27-31
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

The additional search fees were accompanied by the applicant's protest.

No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORTInternational application No.
PCT/US96/01041**BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING**

This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group I, claim(s)1-19, drawn to a nucleic acid molecule encoding a CAL protein, classified in Class 536, subclass 23.6, for example.

Group II, claims 20-22, drawn to a CAL protein, classified in Class 530, subclass 350, for example.

Group III, claims 23-24, drawn to an antibody to a CAL protein, classified in Class 424, subclass 130.1, for example.

Group IV, claim 25, drawn to a truncated CAL protein, classified in Class 530, subclass 300, for example.

Group V, claim 26, drawn to an antibody to a truncated CAL protein, classified in Class 424, subclass 130.1, for example.

Group VI, claim(s) 27-31, drawn to a method of identifying a modified CAL gene which does not encode a protein, classified in Class 435, subclass 6, for example.

The inventions listed as Groups I-VI do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Groups I-V are drawn to a gene encoding a specific CAL protein or a protein having a degree of sequence similarity thereto, while Group VI is drawn to any modified CAL gene which does not encode a functional protein, and to hybridization methods for identifying the gene, wherein the modified non-functional gene and hybridization methods of Group VI are not required by the inventions of Group I-V, and the genes encoding specific proteins of Groups I-V are not required by the invention of Group VI. Furthermore, the inventions of Groups I-III are not linked by a single special technical feature because they are not drawn to a single gene sequence or a single protein sequence, or a single antibody to a single protein sequence. The inventions of Groups I-III are not linked by a single special technical feature to the inventions of Groups IV-V, because the inventions of Groups I-III are not linked by a single sequence, and because the inventions of Groups IV-V involve a truncated protein which is not involved in the inventions of Groups I-III. The inventions of Groups IV and V are not linked by a single special technical feature because they are drawn to the physiologically divergent products of a protein and an antibody, and because Group V is drawn to any of a number of divergent types of antibodies which could bind to the protein of Group IV.